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(A) Humanised antibodies.

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© CDR-grafted antibody heavy and light chains comprise acceptor framework and donor antigan binding regions, the heavy chains comprising donor residues at at least one of positions (6, 23) and/or (24, 48) and/or (48, 71) and/or (73) and/or (73

1 GAATTCCCAA AGACAAAatg gatttteaag tgcagatttt cagcttcctg 51 ctaatcaqtq cctcaqtcat astatecaqa qqacaaattq ttctcaccca 101 gtotocagca atcatgtotg catotocagg ggagaaggto accatgacot 151 gcagtgocag ctcaagtgta agttacatga actggtacca gcagaagtca 201 ggcacctccc ccasaagatg gatttatgac acatecasac tggcttctgg .251 agtcoctgct cacttcaggg gcagtgggtc tgggacctct tactctctca 301 caatcagogg catggaggot gaagatgotg coacttatta otgocagoag 351 tggagtagta accoattoac gttcggctcg gggacaaagt tggaaataas 401 cogggetgat actgcaccaa ctgtatecat cttcccacca tecaqtqaqc 451 agttamcatc tggaggtgcc tcagtcgtgt gcttcttgam camcttctac 501 cccaaagaca tcaatgtcaa gtggaagatt gatggcagtg aacgacaaaa 551 tggcgtcctg aacagttgga ctgatcagga cagcaaagac agcacctaca 601 gcatgagong caccottoacg ttgaccaagg acgagtatga acgacataac 651 agetatacet gtgaggeeac tcacaagaca tcaacttcac ccattgtcaa 701 gagettease aggastgagt gtTAGAGACA AAGGTCCTGA GACGCCACCA 751 CCAGCTCCCA GCTCCATCCT ATCTTCCCTT CTAAGGTCTT GGAGGCTTCC 801 CCACAAGCGC ETACCACTGT TGCGGTGCTC TAAACCTCCT CCCACCTCCT 851 TOTCCTCCTC CTCCCTTTCC TTGGCTTTTA TCATGCTAAT ATTTGCAGAA 

Fig. 1(a)

1 MODEWOITSP LILEARUITS ROOTWINGS AIMSARPORK WITHCOASS 51 VEYNONYOOK SOTSPRWIT DESKLASOVE MIFROSGOT SYSLITSON 101 ANDANTYOO GASSRITTED SOTALIDERA DYARVISTP PESEQLISOO 151 ASVOCTUME YEROINYMWK IDOSERGEN LIMSTOQOK DETFONSSIL 101 TUTNOTERRE HISTCOARMS ETSTEVINES NOMECO

Fig. 1(b)

## Field of the Invention

The present invention relates to humanised antibody molecules, to processes for their production using recombinant DNA technology, and to their therapeutic uses.

The term "humanised antibody molecule" in used to describe a molecule having an antigen binding site derived from an immunoglobulin from a non-human species, and remaining immunoglobulin-derived parts of the molecule being derived from a human immunoglobulin. The antigen binding site typically comprises complementarity determining regions (CDRs) which determine the binding specificity of the antibody molecule and which are carried on appropriate framework regions in the variable domains. There are 3 TO CDRs (CDR), CDR2 and CDR3) in each of the heavy and light chain variable domains.

In the description, reference is made to a number of publications by number. The publications are listed in numerical order at the end of the description.

## Background of the Invention

Natural immunoglobulins have been known for many years, as have the various fragments thereof, such as the Fab. (Fab'p; and Fc fragments, which can be derived by enzymatic cleavage. Natural immunoglobulins comprise a generally Y-shaped molecule having an artigen-blinding site towards the end of each upper arm. The remainder of the structure, and particularly the stem of the Y, mediates the effector of functions associated with immunoglobulins.

Natural immunoplobuline have been used in assay, diagnosis and, to a more limited extent, therapy, However, such uses, especially in therapy, were hindered until recently by the polyclonal nature of natural immunoglobulins. A significant step towards the realisation of the potential of immunoglobulins as therapeutic agents was the discovery of procedures for the production of monoclonal antibodies (MAbs) of defined as specificity (1).

However, most MAbs are produced by hybridomas which are fusions of rodent spleen cells with rodent myeloma cells. They are therefore essentially rodent proteins. There are very few reports of the production of human MAbs.

Since most available MAbs are of rodent origin, they are naturally antigenic in humans and thus can give rise to an undesirable immune response termed the NAMA (Human Anti-Mouse Antibody) response. Therefore, the use of rodent MAbs as therapeutic agents in humans is inherently limited by the fact that the human subject will mount an immunological response to the MAb and will either remove it entirely or at least reduce lits effectiveness. In practice, MAbs of rodent origin may not be used in patients for more than or a rate of the removal tender of the removal tenders of the sevent as well as spling rise to undesirable reactions. For instance, OKT3 a mouse igG2ak MAb which recognises an antigen in the T-cell receptor-CD3 complex has been approved for use in many countries throughout the world as an immunosuppressant in the treatment of such allograft rejection (Chalenoud et all (2) and Jeffers et all (3). I however, in view of the rodent nature of this and other such MAbs, a significant HAMA response which may include a major anti-fully per component, may build up on use. Clearly, it would be highly desirable to diminish or abolish this undesirable HAMA response and thus enlarge the areas of use of these very useful antibiodies.

Proposals have therefore been made to render non-human MAbs less antigenic in humans. Such techniques can be generically termed "humanisation" techniques. These techniques typically involve the use of recombinant DNA technology to manipulate DNA sequences encoding the polypeptide chains of the as mithody molecule.

Early methods for humanising MAbs involved production of chimeric antibodies in which an antigen defining site comprising the complete variable domains of one antibody is linked to constant domains derived from another antibody. Methods for carrying out such chimerisation procedures are described in EP0120894 (Ceitlioch Limited), EP0120824 (Genentech Inc. and City of Hope), EP-A-0 171498 (Res. Dev. Cop. Japan), EP-A-0 173 494 (Stanford University), and W0 8801533 (Ceitlioch Limited). This latter Ceittech application (WO 8801533) discloses a process for proparing an antibody molecule having the variable domains from a mouse MAb and the constant domains from a human immunoglobulin. Such humanized chimeric antibodies, however, still contain a significant proportion of non-human amino acid sequence, i.e. the complete non-human variable domains, and thus may still elicit some HAMA response, services and the service of the servi

In an alternative approach, described in EP-A-0239400 (Winter), the complementarity determining regions (CDRs) of a mouse MAb have been grafted onto the framework regions of the variable domains of unman immunoglobulin by site directed mutagenesis using long origonucleotides. The present invention

relates to humanized antibody molecules prepared according to this alternative approach, i.e. CDR-grafted humanised antibody molecules. Such CDR-grafted humanized antibodies are much less likely to give rise to a HAMA response than humanised chimeric antibodies in view of the much lower proportion of non-human amino acid sequence which they contain.

The earliest work on humanizing MAbs by CDR-grafting was carried out on MAbs recognizing synthetic and a ret MAb recognizing an antigen on human T-cells were humanised by CDR-grafting have been described by Verhoeyen et al. (6) and Riechmann et al. (6) respectively. The preparation of CDR-grafted antibody to the antigen on human T cells is also described in Vol. 980/97482 (Macical Research Council).

In Riechmann et al/Medical Research Council it was found that transfer of the CDR regions alone [as defined by Kabat refs. (7) and (8)] was not sufficient to provide satisfactory antigen inoling activity in the CDR-grafted product. Riechmann et al found that it was necessary to convert a serine residue at position 27 of the human sequence to the corresponding rat phenylalarine residue to obtain a CDR-grafted product having improved antigen binding activity. This residue at position 27 of the heavy chain is within the 1st structural loop adjacent to CDR1. A further construct which additionally contained a human serine to rat tyrosine change at position 39 of the heavy chain is death of the control of th

Very recently Queen et al (3) have described the preparation of a humanised antibody that binds to the interleukin 2 receptor, by combining the CDRs of a murine MAb (anti-Tac) with human immunopiculinin framework and constant regions. The human framework regions were chosen to maximise homology with the anti-Tac MAb sequence. In addition computer modelling was used to Identify framework amino acid residues which wore likely to interact with the CDRs or antigen, and mouse amino acids were used at these positions in the humanised antibody.

In WO 9007981 Gueen et al propose four criteria for designing humanised immunojobulins. The first criterion is to use as the human acceptor the framework from a particular human immunojobulin that is au unusually homologous to the non-human donor immunojobulin to be humanised, or to use a consensus framework from many human antibodies. The second criterion is to use the donor amino acid rather than the acceptor if the human acceptor residue is unusual and the donor residue is typical for human sequences at a specific residue or the framework. The third criterion is to use the donor framework armino acid residue rather than the acceptor at positions immodately adjacent to the CDRs. The fourth criterion is to use the donor framework armino acid residue rather than the acceptor at positions immodately adjacent to the CDRs. The fourth criterion is to use the donor framework armino acid residue of the residue at framework positions at which the amino acid is predicted to have a side chain abom within about 3 Å of the CDRs in a three-dimensional immunojobulin it is proposed that criteria two, thee or four may be applied in addition or alternatively to criterion one, and may be applied upon the companion of the compan

WO 9007881 describes in detail the preparation of a single CDR-grafted humanised antibody, as humanised antibody having specificity for the p55 Tac protein of the IL-2 receptor. The combination of all four criteria, as above, were employed in designing this humanized antibody, the variable region frameworks of the human antibody Eu (7) being used as acceptor. In the resultant humanized antibody the other CDRs were as defined by Kobate et al (7 and 8) and in addition the mouse donor residues were used in the place of the human acceptor residues, at positions 27, 30, 48, 66, 67, 89, 91, 94, 103, 104, 105 and 107 in the heavy chain and a positions 48, 60 and 63 in the light chain, of the variable region frameworks. The humanised arti-Tac antibody obtained is reported to have an affinity for p55 of 3 x 10<sup>9</sup> M<sup>-1</sup>, about one-third of that of the writee MAD.

We have further investigated the proparation of CDR-grafted humanised antibody molecules and have identified a hierarchy of positions within the framework of the variable regions (i.e. outside both the Kabat CDRs and structural loops of the variable regions) at which the amino acid identifies of the residues are important for obtaining CDR-grafted products with satisfactory binding affinity. This has enabled us to establish a protocol for obtaining satisfactory CDR-grafted products which may be applied very widely irrespective of the level of homology between the donor immunoglobulin and acceptor framework. The set of residues which we have identified as being of critical importance does not coincide with the residues identified by Queen et al (9).

## Summary of the Invention

Accordingly, in a first aspect the invention provides a CDR-grafted antibody heavy chain having a variable region domain comprising acceptor framework and donor antigen binding regions wherein the framework comprises donor residues at at least one of positions 6, 23 and/or 24, 48 and/or 49, 71 and/or 73. 75 and/or 76 and/or 78 and 88 and/or 91.

In preferred embodiments, the heavy chain framework comprises donor residues at positions 23, 24, 49, 71, 73 and 78 or at positions 23, 24 and 49. The residues at positions 71, 73 and 78 of the heavy chain framework are preferably either all acceptor or all donor residues.

In particularly preferred embodiments the heavy chain framework additionally comprises donor residues at one, some or all of positions 6, 37, 48 and 94. Also it is particularly preferred that residues at positions of the heavy chain framework which are commonly conserved across species, to positions 2, 4, 25, 36, 39, 47, 33, 103, 104, 106 and 107, if not conserved between donor and acceptor, additionally comprise donor residues. Most preferably the heavy chain framework additionally comprises donor residues at positions 2, 4, 8, 25, 36, 37, 39, 47, 48, 39, 34, 103, 104, 106 and 107.

In addition the heavy chain framework optionally comprises donor residues at one, some or all of positions:

1 and 3,

72 and 76, 20 69 (if 48 is different between donor and acceptor).

69 (if 48 is different between donor and acceptor) 38 and 46 (if 48 is the donor residue).

80 and 20 (if 69 is the donor residue),

67

82 and 18 (if 67 is the donor residue),

25 91,

88. and

any one or more of 9, 11, 41, 87, 108, 110 and 112.

In the first and other aspects of the present invention reference is made to CDR-grafted antibody products comprising acceptor framework and donor antigen binding regions. It will be appreciated that the service widely applicable to the CDR-grafting of antibodies in general. Thus, the donor and acceptor antibodies may be derived from animals of the same species and even same artibody class or sub-class. More usually, however, the donor and acceptor antibodies are derived from animals of different species. Trypically the donor antibody is a non-human antibody, such as a rodent MAb, and the acceptor antibodes and animals of different species.

In the first and other aspects of the present invention, the donor antigen binding region typically comprises at least one CDR from the donor antiboty. Usually the donor antigen binding region comprises at least two and preferably all three CDRs of each of the heavy chain and/or light chain variable regions. The CDRs may comprise the Kabat CDRs, the structural loop CDRs or a composite of the Kabat and structural loop CDRs and any combination of any of these. Preferably, the antigen binding regions of the 40 CDR-grafted heavy chain variable domain comprise CDRs corresponding to the Kabat CDRs at CDRs (residues 50-65) and CDR3 (residues 95-100) and a composite of the Kabat and structural loop CDRs at CDRI (residues 50-35).

The residue designations given above and elsewhere in the present application are numbered according to the Kabat numbering feets. (7) and (8). Thus the residue designations do not always correspond directly with the linear numbering of the amino acid residues. The actual linear amino acid sequence may contain fewer or additional amino acids than in the strict Kabat numbering corresponding to a shortening of, or insertion into, a structural component, whether framework or CDR, of the basic variable domain structure. For example, the heavy chain variable region of the anti-Tac antibody described by Queen et al. (9) contains a single amino acid insert (residue 52a) after residue 52 of CDR2 and a three amino acid insert (residue 52a) after residue 52 of CDR2 and a three amino acid insert (residue 52b) after transevork residue 82, in the Kabat numbering. The correct Kabat numbering of residues may be determined for a given antibody by alignment at regions of homology of the sequence of the antibody with a "standard" Kabat numbered sequence.

The invention also provides in a second aspect a CDR-grafted antibody light chain having a variable region domain comprising acceptor framework and donor antigen binding regions wherein the framework comprises donor residues at at least one of positions 1 and/or 3 and 46 and/or 47. Preferably the CDR grafted light chain of the second aspect comprises donor residues at positions 46 and/or 47.

The invention also provides in a third aspect a CDR-grafted antibody light chain having a variable region domain comprising acceptor framework and donor antigen binding regions wherein the framework com-

prises donor residues at at least one of positions 46, 48, 58 and 71.

In a preferred embodiment of the third aspect, the framework comprises donor residues at all of positions 46, 48, 58 and 71.

In particularly preferred embodiments of the second and third aspects, the framework additionally comprises donor residues at positions 36, 44, 47, 85 and 67. Similarly positions of the light chain framework which are commonly conserved across species, i.e. positions 2, 4, 6, 35, 49, 26, 44-99, 99, 99, 101 and 102, if not conserved between donor and acceptor, additionally comprise donor residues. Most preferably the light chain framework additionally comprises donor residues at positions 2, 4, 6, 35, 36, 38, 44, 47, 49, 26, 64-99, 85, 87, 98, 99, 101 and 102.

In addition the framework of the second or third aspects optionally comprises donor residues at one, some or all of positions:

1 and 3,

33,

60 (if 60 and 54 are able to form at potential saltbridge), 75 70 (if 70 and 24 are able to form a potential saltbridge),

73 and 21 (if 47 is different between donor and acceptor),

37 and 45 (if 47 is different between donor and acceptor), and

any one or more of 10, 12, 40, 80, 103 and 105.

Preferably, the antigen binding regions of the CDR-grafted light chain variable domain comprise CDRs corresponding to the Kabat CDRs at CDR1 (residue 24-34), CDR2 (residues 50-59) and CDR3 (residues 89-97).

The invention further provides in a fourth aspect a CDR-grafted antibody molecule comprising at least one CDR-grafted heavy chain and at least one CDR-grafted light chain according to the first and second or as first and third aspects of the invention.

The humanised artibody molecules and chains of the present invention may comprise: a complete antibody molecule, having full length heavy and light chains; a fragment thereof, such as a Fab, (Fab\*)2 or FV fragment; a light chain or heavy chain moment or dimer; or a single chain antibody, e.g. a single chain FV in which heavy and light chain variable regions are joined by a peptide linker; or any other CDR-grafted on molecule with the same specificity as the original donor artibody. Similarly the CDR-grafted heavy and light chain variable region may be combined with other artibody domains as appropriate.

Also the heavy or light chains or humanised antibody molecules of the present invention may have attached to them an effector or reporter molecule. For instance, it may have a macrocycle, for chelating a heavy metal atom, or a toxin, such as ricin, attached to it by a covalent bridging structure. Alternatively, the sproedures of recombinant DNA technology may be used to produce an immunoglobulin molecule in which the Fc fragment or CH3 domain of a complete immunoglobulin molecule has been replaced by, or has attached thereto by peptide linkage, a functional non-immunoglobulin protein, such as an enzyme or toxin molecule.

Any appropriate acceptor variable region framework sequences may be used having regard to 
colleasybpe of the donor artibody from which the antigen binding regions are derived. Preferably, the type of 
acceptor framework used is of the same/similar class/type as the donor artibody. Conveniently, the 
framework may be chosen to maximise/optimise hornology with the donor artibody sequence particularly at 
positions close or adjacent to the CDHs. However, a high level of hornology between donor and acceptor 
sequences is not important for application of the present invention. The present invention identifies a 
felicarchy of framework residue positions at which donor residues may be important or desirable for 
obtaining a CDR-grated antibody product having satisfactory binding properties. The CDR-grated products 
usually have binding affirities of at least 10° M<sup>-</sup>1, preferably at least about 10° M<sup>-</sup>1, or 
acceptor antibodies irrespective of the level of hornology between their sequences. A protocol for applying 
to the invention to any particular donor-acceptor antibody pair is given hereinafter. Examples of human 
frameworks which may be used are KOL, INVM, REI, EU, LIY and POM (refs. 4 and 5) and the like; for 
instance KOL and NEWM for the heavy chain and REI for the light chain and EU, LAY and POM for both 
he heavy chain and the light for the services.

Also the constant region domains of the products of the invention may be selected having regard to the proposed function of the antibody in particular the effector functions which may be required. For example, the constant region domains may be human IgA, IgE, IgG or IgM domains. In particular, IgG human constant region domains may be used, especially of the IgG1 and IgG3 isotypes, when the humanised antibody molecule is intended for therapoutic uses, and antibody effector functions are recurred. Alter-

natively, IgG2 and IgG4 isotypes may be used when the humanised antibody molecule is intended for therapeutic purposes and antibody effector functions are not required, e.g. for simple blocking of lymphokins activity.

However, the remainder of the artithody molecules need not comprise only protein sequences from immunoglobulins. For instance, a gene may be constructed in which a DNA sequence encoding part of a human immunoglobulin chain is tused to a DNA sequence encoding the amino acid sequence of a functional polyopetide such as an effector or reporter molecule.

Preferably the CDR-grafted antibody heavy and light chain and antibody molecule products are produced by recombinant DNA technology.

Thus in further aspects the invention also includes DNA sequences coding for the CDR-grafted heavy and light chains, cloning and expression vectors containing the DNA sequences, host cells transformed with the DNA sequences and processes for producing the CDR-grafted chains and antibody molecules comprising expressing the DNA sequences in the transformed host cells.

The general methods by which the vectors may be constructed, transfection methods and culture methods are well known per se and form no part of the invention. Such methods are shown, for instance, in references 10 and 11.

The DNA sequences which encode the donor amino acid sequence may be obtained by methods well known in the art. For example the donor coding sequences may be obtained by genomic cloning, or cDNA cloning from suitable hybriddoma cell lines. Positive clones may be screened using appropriate probes for the heavy and light chain genes in question. Also PCR cloning may be used.

DNA coding for acceptor, e.g. human acceptor, sequences may be obtained in any appropriate way. For example DNA sequences coding for preferred human acceptor frameworks such as KOL, REI, EU and NEWM. are widely available to workers in the art.

The standard techniques of molecular biology may be used to prepare DNA sequences coding for the 26 CDF-grafted products. Desired DNA sequences may be synthesised completely or in part using oligonucleotide synthesis techniques. Site-directed mutagenesis and polymerase chain reaction (PCR) techniques may be used as appropriate. For example oligonucleotide directed synthesis as described by Jones et al. (ref. 20) may be used. Also oligonucleotide directed mutageness of a pre-existing variable region as, for example, described by Verhoeyen et al. (ref. 5) or Riectmann et al. (ref. 6) may be used. Also 90 enzymatic filling in of gapped oligonucleotides using T<sub>4</sub> DNA polymerase as, for example, described by Queen et al. (ref. 9) may be used.

Any suitable host cell/vector system may be used for expression of the DNA sequences coding for the CRR-grafted heavy and light chains. Bacteriat o.g. E. coll, and other microbial systems may be used, in particular for expression of antibody fragments such as FAb and (Fab'); fragments, and especially FV fragments and single chain antibody fragments e.g. single chain FVs. Eucaryotic e.g. mammalian host cell expression systems may be used for production of larger CDR-grafted antibody products, including complete antibody molecules. Sultable mammalian host cells include CHO cells and myeloma or hybridoma cell lines.

Thus, in a further aspect the present invention provides a process for producing a CDR-grafted antibody product comprising:

 (a) producing in an expression vector an operon having a DNA sequence which encodes an antibody heavy chain according to the first aspect of the invention;

(b) producing in an expression vector an operon having a DNA sequence which encodes a complementary antibody light chain according to the second or third aspect of the invention;

(c) transfecting a host cell with the or each vector; and

(d) culturing the transfected cell line to produce the CDR-grafted antibody product.

The CDR-grafted product may comprise only heavy or light chain derived polypeptide, in which case only a heavy chain or light chain polypeptide coding sequence is used to transfect the host cells.

For production of products comprising both heavy and light chains, the cell line may be translated with two vectors, the first vector may contain an operon encoding a light chain-derived polypeptide and the second vector containing an operon encoding a heavy chain-derived polypeptide. Preferably, the vectors are identical, except in so far as the coding sequences and selectable markers are concerned, so as to ensure as far as possible that each polypeptide chain is equally expressed. Alternatively, a single vector may be used, the vector including the sequences encoding both light chain- and heavy chain-derived polypeptides.

The DNA in the coding sequences for the light and heavy chains may comprise cDNA or genomic DNA or both. However, it is preferred that the DNA sequence encoding the heavy or light chain comprises at least partially, exen

The present invention is applicable to antibodies of any appropriate specificity. Advantageously, however, the invention may be applied to the humanisation of non-human antibodies which are used for in vivo therapy or diagnosis. Thus the antibodies may be site-specific antibodies such as tumour-specific or cell surface-specific antibodies, suitable for use in in vivo therapy or diagnosis, e.g., tumour imaging. Examples of cell surface-specific antibodies, so are arti-To call antibodies, such as arti-CD3, and CD4 and adhesion molecules, such as CR3, ICAM and ELAM. The antibodies may have specificity for interleutian (including lymphokinus, growth factors and stimulating factors), hormones and other biologically active compounds, and receptors for any of these. For example, the antibodies may have specificity for any of the following: interferons α, β, γ or δ, lt.1, IL2, IL3, or IL4, etc., TNF, GCSF, GMCSF, EPO, IAH, or insulin, etc.

The the present invention also includes therapeutic and diagnostic compositions comprising the CDR-

grafted products of the invention and uses of such compositions in therapy and diagnosis.

Accordingly in a further aspect the invention provides a therapeutic or diagnostic composition comprising a CDR-grafted entibody heavy or light chain or molecule according to previous aspects of the invention in combination with a pharmaceutically accordable center, offluent or excipient.

Accordingly also the invention provides a method of therapy or diagnosis comprising administering an effective amount of a CDR-grafted antibody heavy or light chain or molecule according to previous aspects of the invention to a luman or animal subject.

A preferred protocol for obtaining CDR-grafted antibody heavy and light chains in accordance with the present invention is set out below together with the rationale by which we have derived this protocol. This protocol and rationale are given without prejudice to the generality of the invention as hereinbefore described and defined.

## Protocol

26 It is first of all necessary to sequence the DNA coding for the heavy and light chain variable regions of the donor artibody, to determine their amino acid sequences. It is also necessary to choose appropriate acceptor heavy and light chain variable regions, of known amino acid sequence. The CDR-grafted chain is then designed starting from the basis of the acceptor sequence. It will be appreciated that in some cases the donor and acceptor amino acid residues may be identical at a particular position and thus no change of a acceptor framework residue is required.

 As a first step donor residues are substituted for acceptor residues in the CDRs. For this purpose the CDRs are preferably defined as follows:

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Heavy chain - CDR1: residues 26-35
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- CDR2: residues 50-65
- CDR3: residues 95-102
- Light chain CDR1: residues 24-34
  - CDR2: residues 50-56 - CDR3: residues 89-97
- The positions at which donor residues are to be substituted for acceptor in the framework are then chosen as follows, first of all with respect to the heavy chain and subsequently with respect to the light chain.
- 2. Heavy Chain
- 2.1 Choose donor residues at all of positions 23, 24, 49, 71, 73 and 78 of the heavy chain or all of positions 23, 24 and 49 (71, 73 and 78 are always either all donor or all acceptor).
- 2.2 Check that the following have the same amino acid in donor and acceptor sequences, and if not preferably choose the donor: 2, 4, 6, 25, 36, 37, 39, 47, 48, 93, 94, 103, 104, 106 and 107.
  - 2.3 To further optimise affinity consider choosing donor residues at one, some or any of:
  - i. 1. 3

50

- ii. 72, 76
- iii. If 48 is different between donor and acceptor sequences, consider 69
  - iv. If at 48 the donor residue is chosen, consider 38 and 46 v. If at 69 the donor residue is chosen, consider 80 and then 20
  - d 67
  - vii. If at 67 the donor residue is chosen, consider 82 and then 18
- viii. 91
  - ix. 88 x. 9. 11. 41. 87. 108. 110, 112
- 3. Light Chain

- 3.1 Choose donor at 46, 48, 58 and 71
- 3.2 Check that the following have the same amino acid in donor and acceptor sequences, if not preferably choose donor:
- 2, 4, 6, 35, 38, 44, 47, 49, 62, 64-69 inclusive, 85, 87, 98, 99, 101 and 102
- 3.3 To further optimise affinity consider choosing donor residues at one, some or any of:
  - i. 1, 3 ii. 63
  - iii. 60, if 60 and 54 are able to form potential saltbridge
  - iv. 70, if 70 and 24 are able to form potential saltbridge
  - v. 73, and 21 if 47 is different between donor and acceptor
    - vi. 37, and 45 if 47 is different between donor and acceptor
    - vii. 10, 12, 40, 80, 103, 105

## Rationale

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In order to transfer the binding site of an antibody into a different acceptor framework, a number of factors need to be considered.

## The extent of the CDRs

The CDRs (Complementary Determining Regions) were defined by Wu and Kabat (refs. 4 and 5) on the basis of an analysis of the variability of different regions of antibody variable regions. Three regions per domain were recognised. In the light chain the sequences are 24-34, 50-58, 89-97 (numbering according to Kabat (ref. 4), Eu Index) inclusive and in the heavy chain the sequences are 31-35, 50-85 and 95-102 inclusive.

When antibody structures became available it became apparent that these CDR regions corresponded in the main to loop regions which extended from the 6 barrel framework of the light and heavy variable domains. For H1 there was a discrepancy in that the loop was from 28 to 32 inclusive and for H2 the loop was 52 to 56 and for 12 from 50 to 53. However, with the exception of H1 the CDR regions encompassed the loop regions and extended into the \$\textit{\mu}\$ stand frameworks. In H1 residue 28 tends to be a serine and 27 a phenylalarine or tyrosine, residue 29 is a phenylalarine in most cases. Residues 29 and 30 which are surface residues exposed to selvent might be involved in adigon-binding. A prudent definition of the H1 CDR therefore would include residues 28-35 to include both the loop region and the hypervariable residues 33-35.

It is of interest to note the example of Riechmann et all (ref. 3), who used the residue 31-35 choice for CDR-H1. In order to produce efficient antigen binding, residue 27 also needed to be recruited from the donor (rail) antibody.

### 2. Non-CDR residues which contribute to antigen binding

By examination of available X-ray structures we have identified a number of residues which may have an effect on net artigen binding and which can be demonstrated by experiment. These residues can be sub-divided into a number of groups.

- 2.1 Surface residues near CDR [all numbering as in Kabat et al (ref. 7)].
  - 2.1.1. Heavy Chain Key residues are 23, 71 and 73. Other residues which may contribute to a lesser extent are 1, 3 and 76. Finally 25 is usually conserved but the murine residue should be used if there is a difference.
  - 2.1.2 Light Chain Many residues close to the CDRs, eg. 63, 65, 67 and 68 are conserved. If conserved none of the surface residues in the light chain are likely to have a major effect. However, if the murine residue at these positions is unusual, then it would be of benefit to analyse the likely contribution more obsely. Other residues which may also contribute to binding are 1 and 3, and also 66 of of 70 if the residues at these positions and at 54 and 24 respectively are notentially able to form a sait bridge i.e. 60 + 54, 70 + 24.
- 50 2.2 Packing residues near the CDRs.
  - 2.2.1. Heavy Chain Key residues are 24, 48 and 78. Other key residues would be 36 if not a tryptophan, 94 if not an arginine, 104 and 106 if not glycines and 107 if not a threonine. Residues which may make a further contribution to stable packing of the heavy chain and hence improved affinity are 2, 4, 6, 38, 46, 67 and 59. 67 packs against the CDR residue 63 and this pair could be either both mouse or both human. Finally, residues which contribute to packing in this region but from a longer range are 18, 20, 80, 82 and 86. 82 packs against 67 and in turn 18 packs against 82. 80 packs against 69 and in turn 20 packs against 80. 86 forms an H bond network with 38 and 46. Many of the mouse-human differences appear minor e.g. Leufle, but could have an minor

impact on correct packing which could translate into altered positioning of the CDRs.

2.2.2. Light Chain - Key residues are 48, 58 and 71. Other key residues would be 6 if not glutamine, 35 if not tryptophan, 62 if not phenylalamine or tryosine, 64, 66, 68, 98 and 10 if in of pycines and 102 if not a thronoine. Residues which make a further contribution are 2, 4, 37, 45 and 47. Finally residues 73 and 21 and 19 may make long distance packing contributions of a minor nature.

2.3. Residues at the variable domain interface between heavy and light chains - In both the light and heavy chains most of the non-CDR interface residues are conserved. If a conserved residue is replaced by a residue of different character, e.g. size or charge, it should be considered for retention as the muriner residue.

2.3.1. Heavy Chain - Residues which need to be considered are 37 if the residue is not a valine but is of larger side chain volume or has a charge or polarity. Other residues are 39 if not a glutamine, 64 if not a levoine, 47 if not a typtophan, 91 if not a phenylalamine or tyrosine, 93 if not an alarine and 103 if not a typtophan. Residue 99 is also at the interface but is not in a position where the side chain could be of great impact.

2.3.2. Light Chain - Residues which need to be considered are 36, if not a tyrosine, 38 if not a glutamine, 44 if not a proline, 46, 49 if not a tyrosine, residue 85, residue 87 if not a tyrosine and 88 if not a horwisa

2.4. Variable-Constant region interface - The elbow angle between variable and constant regions may be affected by alterations in packing of key residues in the variable region against the constant region mich may affect the position of V<sub>a</sub> and V<sub>h</sub> with respect to one another. Therefore it is worth noting the residues likely to be in contact with the constant region. In the heavy chain the surface residues potentially in contact with the variable region are conserved between mouse and human arithodies therefore the variable region contact region contact points vary, and the V & C regions are not in such close proximity as the heavy chain. Therefore the influences of the light chain V-C interface may be minor.

2.4.1. Heavy Chain - Contact residues are 7, 11, 41, 87, 108, 110, 112.

2.4.2. Light Chain - In the light chain potentially contacting residues are 10, 12, 40, 80, 83, 103 and 105.

The above analysis coupled with our considerable practical experimental experience in the CDRgrafting of a number of different antibodies have lead us to the protocol given above.

The present invention is now described, by way of example only, with reference to the accompanying

The present invention is now described, by way of example only, will reference to the accompanying Figures 1 - 13.

### Brief Description of the Figures

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- Figure 1 shows DNA and amino acid sequences of the OKT3 light chain;
  - Figure 2 shows DNA and amino acid sequences of the OKT3 heavy chain;
- 40 Figure 3 shows the alignment of the OKT3 light variable region amino acid sequence with that of the light variable region of the human antibody REI;
  - Figure 4 shows the alignment of the OKT3 heavy variable region amino acid sequence with that of the heavy variable region of the human antibody KOL;
- Figure 5 shows the heavy variable region amino acid sequences of OKT3, KOL and various corresponding CBR grafts;
  - Figure 6 shows the light variable region amino acid sequences of OKT3, REI and various corresponding CDR grafts;
    - Figure 7 shows a graph of binding assay results for various grafted OKT3 antibodies'
    - Figure 8 shows a graph of blocking assay results for various grafted OKT3 antibodies;
- Figure 9 shows a similar graph of blocking assay results;
  - Figure 10 shows similar graphs for both binding assay and blocking assay results;
  - Figure 11 shows further similar graphs for both binding assay and blocking assay results;
  - Figure 12 shows a graph of competition assay results for a minimally grafted OKT3 antibody
- compared with the OKT3 murine reference standard, and shows a similar graph of competition assay results comparing a fully grafted OKT3
  - antibody with the murine reference standard.

## DETAILED DESCRIPTION OF EMBODIMENTS OF THE INVENTION

#### EXAMPLE 1

## 5 CDR-GRAFTING OF OKT3

### MATERIAL AND METHODS

#### 1. INCOMING CELLS

Hybridoma cells producing antibody OKT3 were provided by Ortho (seedlot 4882.1) and were grown up in antibiotic free Dulbecco's Modified Eagles Medium (DMEM) supplemented with glutamine and 5% foetal calf serum, and divided to provide both an overgrown supernatant for evaluation and cells for extraction of RNA. The overgrown supernatant was shown to contain 250 ug/ml. murine IgG2a/kappa

antibody. The supernatant was negative for murine lambda light chain and IgG1, IgG2b, IgG3, IgA and IgM heavy chain, 20mL of supernatant was assayed to confirm that the antibody present was OKT3.

## 2. MOLECULAR BIOLOGY PROCEDURES

Basic molecular biology procedures were as described in Manistis et al (ref. 9) with, in some cases, minor modifications. DRA sequencing was performed as described in Sanger et al (ref. 11) and the Amersham International PIc sequencing handblook. Site directed mutagenesis was as described in Kramer et al (ref. 12) and the Anglian Biotechnology Ltd. handbook. COS cell expression and metabolic labelling studies were as described in Whittee et al (ref. 13)

### 3. RESEARCH ASSAYS

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3.1. ASSEMBLY ASSAYS Assembly assays were performed on supernatants from transfected COS cells to determine the amount of intact IgG present.

3.1.1. COS CELLS TRANSFECTED WITH MOUSE OKT3 GENES The assembly assay for Intact mouse IgG in COS cell supernatants was an ELISA with the following format:

98 well microtitre plates were coated with F(ab)2 goat anti-mouse IgG Fc. The plates were washed in water and samples added for 1 hour at room temperature. The plates were washed and F(ab)2 goat anti-mouse IgG F(ab)2 (HRPO conjugated) was then added. Substrate was added to reveal the reaction. UPC10, a mouse IgG2a myeloma, was used as a standard.

3.12. COS AND CHO CELLS TRANSFECTED WITH CHIMERIC OR CDR-GRAFTED OKT3 GENES

The assembly assay for chimeric or CDR-grafted antibody in COS cell supernatants was an ELISA with the following format:

96 well microtitre plates were coated with F(ab1/2 goat anti-human IgG Fc. The plates were washed and samples added and incubated for 1 hour at room temperature. The plates were washed and monoclonal mouse anti-human kappa chain was added for 1 hour at room temperature.

The plates were washed and Frabh2 goat anti-mouse IgG Fc (HRPC conlugated) was added. Enzyme substrate was added to reveal the maction. Chimaric RP23 (IgG4) (en. 13) was used as a standard. The use of a monoclonal anti-kappa chain in this assay allows grafted antibodies to be and from the chimeric standard.

#### 3.2. ASSAY FOR ANTIGEN BINDING ACTIVITY

Material from COS cell supernatants was assayed for OKT3 antigen binding activity onto CD3 positive cells in a direct assay. The procedure was as follows:

HUT 78 cells (human T cell line, CD3 positive) were maintained in culture. Monolayers of HUT 78 cells were prepared onto 96 well ELISA plates using polyt-t-lysine and glutaraldehyde. Samples were added to the monolayers for 1 hour at room temperature.

The plates were washed gently using PBS. F(ab')2 goat anti-human IgG Fc (HRPO conjugated) or F-(ab')2 goat anti-mouse IgG Fc (HRPO conjugated) was added as appropriate for humanised or mouse sameles. Substrate was added to reveal the reaction.

The negative control for the cell-based assay was chimeric B72.3. The positive control was mouse Orthomune OKT3 or chimeric OKT3, when available. This cell-based assay was difficult to perform, and an alternative assay was developed for CDR-grafted OKT3 which was more sensitive and easier

to carry out.

In this system CDR-grafted OKT3 produced by COS cells was tested for its ability to bind to the CD3-positive HPB-ALL (fluman peripheral blood acute lymphocytic leukemis) cell line. It was also tested for its ability to block the binding or flumine OKT3 to these cells. Binding was measured by the following

procedure: HPB-ALL cells were harvested from tissue culture. Cells were incubated at 4°C for 1 hour with various dilutions of test antibody, positive control antibody, or negative control antibody. The cells were washed once and incubated at 4°C for 1 hour with an FTTC-labelled goat anti-human IgG (Fcspecific, mouse absorbed). The cells were washed twice and analysed by cytofluorography. Chimeric OKT3 was used as a positive control for direct binding. Cells incubated with mock- transfected COS cell supernatant, followed by the FITC-labelled goat anti-human IgG, provided the negative control. To test the ability of CDR-grafted OKT3 to block murine OKT3 binding, the HPB-ALL cells were incubated at 4°C for 1 hour with various dilutions of test antibody or control antibody. A fixed saturating amount of FITC OKT3 was added. The samples were incubated for 1 hour at 4°C, washed twice and analysed by cytofluorography. FTTC-labelled OKT3 was used as a positive control to determine maximum binding. Unlabelled murine OKT3 served as a reference standard for blocking. Negative controls were unstained cells with or without mock-transfected cell supernatant. The ability of the CDR-grafted OKT3 light chain to bind CD3-positive cells and block the binding of murine OKT3 was initially tested in combination with the chimeric OKT3 heavy chain. The chimeric OKT3 heavy chain is composed of the murine OKT3 variable region and the human IgG4 constant region. The chimeric heavy chain gene is expressed in the same expression vector used for the CDR-grafted genes. The CDR-grafted light chain expression vector and the chimeric heavy chain expression vector were co-transfected into COS cells. The fully chimeric OKT3 antibody (chimeric light chain and chimeric heavy chain) was found to be fully capable of binding to CD3 positive cells and blocking the hinding of murine OKT3 to these cells.

3.3 DETERMINATION OF RELATIVE BINDING AFFINITY

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3.3 DETERMINATION OF HELATIVE BIRUNICA AFFIRITY

The relative binding affirities or CDP-grafted anti-CD3 monoclonal antibodies were determined by competition binding affirity or CDP-grafted anti-CD3 monoclonal antibodies were determined by competition binding affirity as a secure of CD3 antigen, and throrsecoin-conjugated murino (KOT3 (FloxATS) of known binding affirity as a tarcer antibody. The binding affirity of Fl-OKT3 tracer antibody was determined by a direct binding assay in which increasing amounts of Fl-OKT3 were incubated with HPP-B-LL (Sct/Pf) in PBS with 5% footal calf sarum for 60 min, at 4°C. Cells were washed, and the fluorescence intensity was determined on a FACScan flow cytometer calibrated with quantitative microbead standards (Flow Cytometry Standards, Research Triangle Park, HO, Fluorescence intensity per antibody molecule (FP ratio) was determined by using microbeads which have a predetermined number of mouse IgG antibody binding sites (Simply Celluter beads, Flow Cytometry Standards). Fire equals the fluorescence intensity of beads saturated with Fl-OKT3 divided by the number of binding sites per bead. The amount of bound and free Fl-OKT3 was calculated from the mean fluorescence intensity per cell, and the ratio of bound/free was plotted against the number of moles of antibody bound. A linear fit was used to determine the affiling of binding desolute value of the slope.)

For competitive binding, increasing amounts of competitor antibody were added to a sub-saturating dose of Fl-OKT3 and incubated with 5x0° HPB-ALL in 200 mil of PBS with 5% foetal calf serum, for 0 min at 4°C. The fluoresconce intensities of the cells were measured on a FACScan flow cytometer calibrated with quantitative microbead standards. The concentrations of bound and free Fl-OKT3 were calculated. The affinities of competing anti-bodies were calculated from the equation DKI-JOKT3] = (I/Kx) - (I/Ka), where Ka is the affinity of murine OKT3, Kx is the affinity of competitor X, [] is the concentration of competitor antibody at which bound/free binding is R/2, and R is the maximal bound/free binding.

## 4. cDNA LIBRARY CONSTRUCTION

4.1. mRNA PREPARATION AND CDNA SYNTHESIS

OKT3 producing cells were grown as described above and 1.2 x 10<sup>9</sup> cells harvested and mRNA extracted using the guaridinium/LIC extraction procedure. cDNA was prepared by priming from OligodT to generate full length cDNA. The cDNA was methylated and EcoR1 linkers added for cloning. 4.2 LIBRARY CONSTRUCTION

The cDNA library was ligated to pSP65 vector DNA which had been EcoR1 cut and the 5' phosphate groups removed by calf intestinal phosphatase (EcoR1/CIP). The ligation was used to transform high transformation efficiency Escherichia coil (E.coli) HB101. A cDNA library was prepared. 3600 colonies were screened for the light chain and 10000 colonies were screened for the heavy chain. 5. SCREENING.

Cool colonies positive for either heavy or light chain probes were identified by oligonucleotide screening using the oligonucleotides: S' TCCAGATGITTAACTGCTCAC for the light chain, which is complementary to a sequence in the mouse kappa constant region, and 5' CAGGGGCCAGTGGATGGATAGAC for the heavy chain which is complementary to a sequence in the mouse ligida constant CHI domain region. 12

light chain and 9 heavy chain clones were identified and taken for second round screening. Positive clones from the second round of screening were grown up and DNA prepared. The sizes of the gene inserts were estimated by gel electrophresis and inserts of a size capable of containing a full length cDNA were subcloned into MI3 for DNA sequencing.

6. DNA SEQUENCING

Clones representing four size classes for both heavy and light chains were obtained in M13. DNA sequence for the 5' untranslated regions, signal sequences, variable regions and 3' untranslated regions of full length cDNAs [Figures 16] and 2(a)] were obtained and the corresponding amino acid sequences predicted [(Figures 16) and 2(b)]. In Figure 1(a) the untranslated DNA regions are shown in uppercase, and in both Ficures 1 and 2 the signal sequences are underfined.

7. CONSTRUCTION OF CDNA EXPRESSION VECTORS

Celltech expression vectors are based on the plasmid pEERicCMV (ref. 14). A polylinker for the insortion of genes to be expressed has been introduced after the major immediate early promoter/enhancer of the human Cytomogalovinus (nCMV). Marker genes for selection of the plasmid in transfected eukaryotic cells can be inserted as BarnH1 cassettes in the unique BarnH1 site of pEEB nCMV; for instance, the own marker to provide pEEB nCMV not. It is usual practice to insert the new and gpt markers prior to insertion of the gene of inforest, whereas the GS marker is inserted last because of the presence of interest is the inthe research.

The selectable markers are expressed from the SV40 late promoter which also provides an origin of replication so that the vectors can be used for expression in the COS cell transient expression system.

The mouse sequences were excised from the M13 based vectors described above as EcoR1 fragments and cloned into either pEE6+hCMV-neo for the heavy chain and into EE6+hCMV-gpt for the light chain to yield vectors pA136 and pA136 respectively.

B. EXPRESSION OF CDNAS IN COS CELLS

Plasmids pJA135 and pJA136 were co-transfected into COS cells and supernatant from the transient expression experiment was shown to contain assembled antibody which bound to T-cell enriched lymphocytes. Metabolic labelling experiments using <sup>55</sup>S methionine showed expression and assembly of heavy and light chains.

9. CONSTRUCTION OF CHIMERIC GENES

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- Construction of chimeric genes followed a previously described strategy (Mhittle et al (ed. 13)). A nestriction site near the 3' end of the variable domain sequence is identified and used to attach an oligonucleotide adapter coding for the remainder of the mouse variable region and a suitable restriction site for attachment to the constant region of choice.
  - 9.1. LIGHT CHAIN GENE CONSTRUCTION
  - The mouse light chain cDNA sequence contains an Aval site near the 3' end of the variable region [Fig. 1(a)]. The majority of the sequence of the variable region was isolated as a 396 bp. ExOR1-Aval fragment. An oligonuclootide adapter was designed to replace the emailated of the 3' region of the variable region from the Aval site and to include the 5' residues of the human constant region up to and including a unique Nar1 site which had been previously engineered into the constant region.
- A Hind111 site was introduced to act as a marker for insertion of the linker.

  The linker was ligated to the V<sub>L</sub> fragment and the 413 bp EcoR1-Nar1 adapted fragment was purified
  - from the ligation mixture.

    The constant region was isolated as an Nart-BamH1 fragment from an M13 clone NW361 and was
  - ligated with the variable region DNA into an EcoR1/BamH1/CIP pSP65 treated vector in a three way reaction to yield plasmid JA143. Clones were isolated after transformation into E.coli and the linker and unction sequences were confirmed by the presence of the Hind111 site and by DNA sequencing. 9.2 LIGHT CHAIN GENE CONSTRUCTION - VERSION 2

The construction of the first chimeric light chain gene produces a fusion of mouse and human amino acid sequences at the variable-constant region junction. In the case of the OKT3 light chain the amino acids at the chimera junction are:

.....Leu-Glu-Ile-<u>Asn-Arq/ -/Thr</u>-Val-Ala -Al

VARIABLE CONSTANT

This arrangement of sequence introduces a potential site for Asparagine (Asn) linked (N-linked) glycosylation at the V-C junction. Therefore, a second version of the chimenic light chain oligonucleotide adapter was designed in which the threonine (Irrh), the first amino acid of the human

constant region, was replaced with the equivalent amino acid from the mouse constant region, Alanine (Ala).

An internal Hind111 site was not included in this adapter, to differentiate the two chimeric light chain

The variable region fragment was isolated as a 376 bp EcoR1-Aval fragment. The oligonucleotide linker was ligated to Narf cut pMW861 and then the adapted 396bp constant region was isolated after recutting the modified pMW361 with EcoR1. The variable region fragment and the modified constant region fragment were ligated directly into EcoR1/CIP treated pEEB/CMVneo to yield pJA137. Initially all clones examined had the insert in the incorrect crientation. Therefore, the insert was re-isolated and reckoned to turn the Insert round and yield plasmid pJA141. Several clones with the insert in the correct orientation were obtained and the adapter sequence of one was confirmed by DNA sequencing

9.3. HEAVY CHAIN GENE CONSTRUCTION

9.3.1, CHOICE OF HEAVY CHAIN GENE ISOTYPE

The constant region isotype chosen for the heavy chain was human IgG4.

9.3.2. GENE CONSTRUCTION

The heavy chain cDNA sequence showed a Ban1 site near the 3' end of the variable region [Fig. 2(a)-]. The majority of the sequence of the variable region was isolated as a 428bp. ExoPIACIPDEan1 fragment. An oligonucleotide adapter was designated to replace the remainder of the 3' region of the variable region from the Ban1 site up to and including a unique Hindfli site which had been previously engineered into the first two amino acids of the constant region.

The linker was ligated to the V<sub>k</sub> fragment and the EcoR1-Hnd111 adapted fragment was purified from the ligation mixture. The variable region was ligated to the constant region by cutting pJA91 with EcoR1 and Hind111 removing the intron fragment and replacing it with the V<sub>k</sub> to yield pJA142. Clores were isolated after transformation into EcoII JM101 and the linker and junction sequences were confirmed by DMA sequencing. (N.E. The Hind111 sits list lost on cloring).

# 10. CONSTRUCTION OF CHIMERIC EXPRESSION VECTORS

10.1. neo AND gpt VECTORS

The chimeric light chain (version 1) was removed from pJA143 as an EcoR1 fragment and cloned into EcoR1/CIP treated pEE6hCM/nee expression vector to yield pJA145. Clones with the insert in the correct orientation were identified by restriction mapping.

The chimeric light chain (version 2) was constructed as described above.

The chimeric heavy chain gene was isolated from pJA142 as a 2.5Kbp EcoR1/BamH1 fragment and cloned into the EcoR1/Bc11/C1P treated vector fragment of a derivative of pEE6hCMVgpt to yield plasmid pJA144.

10.2. GS SEPARATE VECTORS

GS versions of pJA141 and pJA14 were constructed by replacing the nee and gpt cassettes by a versions to PJA17 the plasmids, isolation of the vector fragment and ligation to a GS containing fragment from the plasmid pRO49 to yield the light chain vector pJA179 and the heavy chain vector pJA180.

10.3. GS SINGLE VECTOR CONSTRUCTION

Single vector constructions containing the ct. (chimeric light), ctl (chimeric heavy) and GS genes on one plasmid in the order ct.-ctl-GS, or ctl-ct-GS and with transcription of the genes being head to tail e.g. ct.-ctl-GS were constructed. These plasmids were made by treating plu/179 or plu/189 with BamH1/CIP and ligating in a Bg11/H/ind111 hCMV promoter casestic along with either the Hind111/BamH1 fragment from pJA141 into pJA180 to give the ctl-ct-GS plasmid pJA182 or the Hind111/BamH1 fragment from pJA144 into pJA179 to give the ctl-ctl-GS plasmid pJA181.

11. EXPRESSION OF CHIMERIC GENES

11.1. EXPRESSION IN COS CELLS

11.1. EVPHESSION IN COS CELLS
The chimeric ambitody plasmid pJA145 (ct.) and pJA144 (cfl) were co-transfected into COS cells and supernatent from the transient expression experiment was shown to contain assembled antibody which bound to the HUT 78 human T-cell line. Metabloc liabeling experiments using \*S methiorine showed expression and assembly of heavy and light chains. However the light chain mobility seen on COS cells in the presence of tunicamycin showed a reduction in size of the light chain to that shown for control chimeric antibodies and the OKT3 mouse light chain. Therefore JA141 was constructed and expressed. In this case the light that for the van abernat mobility or a size shift in the presence or absence of tunicamycin. This second version of the chimeric light chain, when expressed in association with chimeric heavy (ctl) chain, conduced antibody which showed good binding to HUT

78 cells. In both cases antigen binding was equivalent to that of the mouse antibody.

11.2 EXPRESSION IN CHINESE HAMSTER OVARY (CHO) CELLS Stable cell lines have been prepared from plasmids PJA141/pJA144 and from pJA179/pJA180, pJA181 and pJA182 by transfection into CHO cells

12. CDR-GRAFTING

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The approach taken was to try to introduce sufficient mouse residues into a human variable region framework to generate antigen binding activity comparable to the mouse and chimeric antibodies.

12.1. VARIABLE REGION ANALYSIS

From an examination of a small database of structures of antibodies and antigen-antibody complexes it is clear that only a small number of antibody residues make direct contact with antigen. Other residues may contribute to antigen binding by positioning the contact residues in favourable configurations and also by inducing a stable packing of the individual variable domains and stable interaction of the light and heavy chain variable domains.

The residues chosen for transfer can be identified in a number of ways:

(a) By examination of antibody X-ray crystal structures the antigen binding surface can be predominantly located on a series of loops, three per domain, which extend from the B-barret temperature.

(b) By analysis of antibody variable domain sequences regions of hypervariability (termed the Complementarity Determining Regions (CDPs) by Wu and Kabat (ref. 6)] can be identified in the most but not all cases these CDRs correspond to, but extend a short way beyond, the loop regions

(c) Residues not identified by (a) and (b) may contribute to antigen binding directly or indirectly by affecting antigen binding site topology, or by inducing a stable packing of the individual variable domains and stabilising the intervariable domain interaction. These residues may be identified either by superimposing the sequences for a given antibody on a known structure and looking at key residues for their contribution, or by sequence alignment analysis and noting "idlosyncratic" residues followed by examination of their structural location and likely effects.

12.1.1. LIGHT CHAIN

Figure 3 shows an alignment of sequences for the human framework region REI and the OKT3 light variable region. The structural loops (LOOP) and CDRs (KABAT) believed to correspond to the antigen binding region are marked. Also marked are a number of other residues which may also contribute to antigen binding as described in 13.1(e). Above the sequence in Figure 3 the residue type indicates the spatial location of each residue side chain, derived by examination of resolved structures from X-ray crystallography analysis. The key to this residue type designation is as follows:

N - near to CDR (From X-ray Structures)

P - Packing B - Buried Non-Packing

S - Surface E - Exposed

I - Interface \* - Interface - Packing/Part Exposed

? - Non-CDR Residues which may require to be left as Mouse sequence. Residues underfined in Figure 3 are amino acids. REI was chosen as the human framework because the light chain is a kappa chain and the kappa variable regions show higher homology with the mouse sequences than a lambda light variable region, e.g. KOL (see below). REI was chosen in preference to another

kappa light chain because the X-ray structure of the light chain has been determined so that a structural examination of individual residues could be made.

12.1.2. HEAVY CHAIN

Kimilarly Figure 4 shows an alignment of soquences for the human framework region KOL and the OKT3 heavy variable region. The structural loops and CDT8 believed to correspond to the antigen binding region are marked. Also marked are a number of other residues which may also contribute to antigen binding as described in 12.1(e). The residue type key and other indicators used in Figure 4 are the same as those used in Figure 3. KOL was chosen as the heavy chain framework because the X-ray structure has been determined to a botter resolution than, for example, NEWM and also the sequence alignment of OKT3 heavy variable region showed a slightly better homology to KOL than to NEWM.

12.2. DESIGN OF VARIABLE GENES

The variable region domains were designed with mouse variable region optimal codon usage [Grantham and Perrin (ref. 15)] and used the B72.3 signal sequences [Whittle et al (ref. 13)]. The

sequences were designed to be attached to the constant region in the same way as for the chimeric genes described above. Some constructs contained the "Kozak consensus sequence" [Kozak (ref. 16)] directly linked to the 5' of the signal sequence in the gene. This sequence motif is believed to have a beneficial role in translation initiation in eukaryotes.

5 12.3. GENE CONSTRUCTION

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To build the variable regions, various strategies are available. The sequence may be assembled by using oligonucleotides in a manner similar to Jones et al. (ef. 17) or by simultaneously replacing like ODRs or loop regions by oligonucleotide directed site specific mutagenesis in a manner similar to Verhooyen et al. (ref. 2). Both strategies were used and a list of constructions is set until nables 1 and 2 and Figures 4 and 5. It was noted in several cases that the mutagenesis approach led to deletions and rearrangements in the gene being remodelled, while the success of the assembly approach was very sensitive to the quality of the oligonucleotides.

13. CONSTRUCTION OF EXPRESSION VECTORS

Genes were isolated from M13 or SP65 based intermediate vectors and cloned into pEE8hCMVneo for the light chains and pEE8hCMVgpt for the heavy chains in a manner similar to that for the chimeric genes as described above.

	TABLE 1		CDR-GRA	FTED C	ENE CO	STRUCTS				
	CODE	HOUSE :	EQUENCE				METHOD OF	KOZAK		
		CONTEN	r				CONSTRUCT	ION	SEQUE	ENCE
5									- 4	
				• • • • •						
	LIGHT C	HAIN	ALL HUM	AN FRA	HEWORK	RE1				
	121	26-32.	50-56,	91-96	inclus	ive	SDM and ge	ene assembly	+	n.d.
10	121A	26-32,	50-56,	91-96	inclus	ive	Partial g	ene assembly	n.d.	+
		+1, 3,	46, 47							
	121B	26-32,	50-56,	91-96	inclus	íve	Partial g	ene assembly	n.d.	+
		+ 46.	47							
15	221	24-24.	50-56.	91-96	inclus	ive	Partial g	ene assembly	+	+
	221A	24-34.	50-56,	91-96	inclus	ive	Partial g	ene assembly	+	+
		+1, 3,	46. 47							
	221B	24-34.	50-56,	91-96	inclus	ive	Partial g	ene assembly	+	+
20		+1, 3								
	221C	24-34.	50-56,	91-96	inclus	ive	Partial g	ene assembly	+	+
25	HEAVY C	HAIN	ALL HU	HAN F	RAMEWOR	K KOL				
	121	26-32,	50-56,	95-10	OB incl	usive	Gene asse	mbly	n.d.	+
	131	26-32,	50-58,	95-10	OB incl	usive	Gene asse	mbly	n.d.	+
	141	26-32,	50-65,	95-10	OB incl	usive	Partial g	ene assembly	+	n.d.
30	321	26-35,	50-56,	95-10	OB incl	usive	Partial g	ene assembly	+	n.d.
	331	26-35	50-58,	95-10	OB incl	usive	Partial g	ene assembly	٠	
							Gene asse	mbly		+
	341	26-35.	50-65.	95-10	OB incl	usive	SDM		+	
35							Partial g	ene assembly		+ .
	341A	26-35	50-65,	95-10	OB incl	usive	Gene asse	mbly	n.d.	+
		+6, 2	. 24, 41	8, 49,	71, 73	, 76,				
		78, 88	. 91 (+	63 <b>-</b> h	uman)					
40	341B	26-35	50-65,	95-10	OB incl	usive	Gene asse	ably	n.d.	+
		+ 48.	49, 71,	73. 7	6, 78,	88. 91				
		(+63 -	human)							
45	Partia	1 gene	Variabl	rected e regi e regi	on asse	embled en	combinati	om oligonucleo	tion	
50	asse	mbly	and gen	e asso iable cs fro	mbly o	r by olig	onucleotic	ginally create ie assembly of n with restric created by SI	part tion	of

## 14. EXPRESSION OF CDR-GRAFTED GENES

14.1. PRODUCTION OF ANTIBODY CONSISTING OF GRAFTED LIGHT (gL) CHAINS WITH MOUSE HEAVY (mH) OR CHIMERIC HEAVY (cH) CHAINS

All gL chains, in association with mH or cH produced reasonable amounts of antibody. Insertion of the Kozak consensus sequence at a position 5' to the ATG (kgl. constructs) however, led to a 2-5 fold

improvement in net expression. Over an extended series of experiments expression levels were raised from approximately 200ng/ml to approximately 500 ng/ml for kgL/cH or kgL/mH combinations.

When direct binding be antigen on HUT 78 cells was measured, a construct designed to include mouse sequence based on loop length (gl.121) did not tead to active antibody in association with mH or cH. A construct designed to include mouse sequence based on Kabat CDRs (gl.221) demonstrated some weak binding in association with mH or cH. However, when framework residues 1, 3, 46, 47 were changed from the human to the murino OKT3 equivalents based on the arguments outlined in Sociation 12.1 antigen binding was demonstrated when both of the new constructs, which were termed 211A and 221A were co-expressed with cH. When the effects of these residues were examined in more detail, it appears that residues 1 and 3 are not major contributing residues as the product of the gl.221B gene shows title describate binding activity in association with cH. The fight thain product of gl.221C, in which mouse sequences are present at 48 and 47, shows good binding activity in association with cH.

14.2 PRODUCTION OF ANTIBODY CONSISTING OF GRAFTED HEAVY (gH) CHAINS WITH MOUSE LIGHT (mL) OR CHIMERIC LIGHT (cL) CHAINS

Expression of the gH genes proved to be more difficult to achieve than for gL. First, inclusion of the Kozak sequence appeared to have no marked effect on expression of gH geness. Expression appears to be slightly improved but not to the same degree as seen for the grafted light chain.

Also, it proved difficult to demonstrate production of expected quantities of material when the loop choice (amino acid 26-32) for CDR1 is used, e.g. gH121, 131, 141 and no conclusions can be drawn about these constructs.

Moreover, co-expression of the gH341 gene with cL or mL has been variable and has tended to produce lower amounts of antibody than the cH/cL or mH/mL combinations. The alterations to gH341 to produce of 4341 A and off-3418 lead to improved levels of expression.

This may be due either to a general increase in the fraction of mouse sequence in the variable region, or to the alteration at position 63 where the residue is returned to the human simino acid Valene (Val) from Phenylatanine (Phe) to avoid possible internal packing problems with the rest of the human framework. This arrandement also occurs in of 1831 and olf3521.

When gH321 or gH331 were expressed in association with ct., antibody was produced but antibody binding activity was not detected.

When the more conservative gH341 gene was used antigen binding could be detected in association with ct. or mt., but the activity was only marginally above the background level. When further mouse residues were substituted based on the arguments in 121, antigen binding could be clearly demonstrated for the antibody produced when kgH341A and kgH341B were expressed in association

with cl.

14.3 PRODUCTION OF FULLY CDR-GRAFTED ANTIBODY

The kgL221A gene was co-expressed with kgH341, kgH341A or kgH341B. For the combination kgH221A/kgH341 very little material was produced in a normal COS cell expression.

For the combinations kgL221A/kgH341A or kgH221A/kgH341B amounts of antibody similar to gL/cH was produced.

In several experiments no antigen binding activity could be detected with kgH221A/gH341 or kgH221A/kgH341 combinations, atthough expression levels were very low.

Antigen binding was detected when kgL221A/kgH341A or kgH221A/kgH341B combinations were expressed. In the case of the antibody produced from the kgL221A/kgH341A combination the antigen binding was very similar to that of the chimeric antibody.

An analysis of the above results is given below.

15. DISCUSSION OF COR-GRAFTING RESULTS In the design of the fully humanised antibody the aim was to transfer the minimum number of mouse amino acids that would confire antiloen binding onto a human antibody framework.

15.1. LIGHT CHAIN

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15.1.1. EXTENT OF THE CDRs

For the light chain the regions defining the loops known from structural studies of other antibodies to contain the antigen contacting residues, and those hypervariable sequences defined by klotal et al (refs. 4 and 5) as Complementally Determining Regions (CDRs) are equivalent for CDR2. For CDR1 the hypervariable region extends from residues 24-34 inclusive while the structural loop extends from 26-25 inclusives. In the case of OKTs there is only one amino acid difference between the two options, at amino acid 24, where the mouse sequence is a serine and the human framework REI has glutamine. For CDR3 the loop extends from residues 34-96 inclusive while the

Kabat hypervariability extends from residues 89-97 inclusive. For OKT3 amino acids 89, 90 and 97 are the same between OKT3 and RE1 (Fig. 3). When constructs based on the loop choice for CDR1 (gL121) and the Kabat choice (gL221) were made and co-expressed with mH or cH no evidence for antigen binding activity could be found for gL121, but trace activity could be detected for the gL221, suggesting that a single extra mouse residue in the grafted variable region could have some detectable effect. Both gene constructs were reasonably well expressed in the transient expression system.

15.1.2. FRAMEWORK RESIDUES

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The remaining framework residues were then further examined, in particular amino acids known from X-ray analysis of other antibodies to be close to the CDRs and also those amino acids which in OKT3 showed differences from the consensus framework for the mouse subgroup (subgroup VI) to which OKT3 shows most homology. Four positions 1, 3, 46 and 47 were identified and their possible contribution was examined by substituting the mouse amino acid for the human amino acid at each position. Therefore gL221A (gL221 + D1Q, Q3V, L46R, L47W, see Figure 3 and Table 1) was made, cloned in EE6hCMVneo and co-expressed with cH (pJA144). The resultant antibody was well expressed and showed good binding activity. When the related genes gL221B (qL221 + D1Q, Q3V) and gL221C (gL221 + L46R, L47W) were made and similarly tested, while both genes produced antibody when co-expressed with cH, only the gL221C/cH combination showed good antigen binding. When the gL121A (gL121 + D1Q, Q3V, L46R, L47W) gene was made and co-expressed with cH, antibody was produced which also bound to antigen.

15.2. HEAVY CHAIN

15.2.1, EXTENT OF THE CDRs For the heavy chain the loop and hypervariability analyses agree only in CDR3. For CDR1 the loop region extends from residues 26-32 inclusive whereas the Kabat CDR extends from residues 31-35 inclusive. For CDR2 the loop region is from 50-58 inclusive while the hypervariable region covers amino acids 50-65 inclusive. Therefore humanised heavy chains were constructed using the framework from antibody KOL and with various combinations of these CDR choices, including a shorter choice for CDR2 of 50-56 inclusive as there was some uncertainty as to the definition of the end point for the CDR2 loop around residues 56 to 58. The genes were co-expressed with mL or cL initially. In the case of the gH genes with loop choices for CDR1 e.g. gH121, gH131, gH141 very little antibody was produced in the culture supernatants. As no free light chain was detected it was presumed that the antibody was being made and assembled inside the cell but that the heavy chain was aberrant in some way, possibly incorrectly folded, and therefore the antibody was being degraded internally. In some experiments trace amounts of antibody could be detected in 35 S labelling studies.

As no net antibody was produced, analysis of these constructs was not pursued further.

When, however, a combination of the loop choice and the Kabat choice for CDR1 was tested (mouse amino acids 26-35 inclusive) and in which residues 31 (Ser to Arg), 33 (Ala to Thr), and 35 (Tyr to His) were changed from the human residues to the mouse residue and compared to the first series, antibody was produced for gH321, kgH331 and kgH341 when co-expressed with cL. Expression was generally low and could not be markedly improved by the insertion of the Kozak consensus sequence 5' to the ATG of the signal sequence of the gene, as distinct from the case of the gL genes where such insertion led to a 2-5 fold increase in net antibody production. However, only in the case of gH341/mL or kgH341/cL could marginal antigen binding activity be demonstrated. When the kgH341 gene was co-expressed with kgL221A, the net yield of antibody was too low to give a signal above the background level in the antigen binding assay.

15.2.2. FRAMEWORK RESIDUES

As in the case of the light chain the heavy chain frameworks were re-examined. Possibly because of the lower initial homology between the mouse and human heavy variable domains compared to the light chains, more amino acid positions proved to be of interest. Two genes kgH341A and kgH341B were constructed, with 11 or 8 human residues respectively substituted by mouse residues compared to gH341, and with the CDR2 residue 63 returned to the human amino acid potentially to improve domain packing. Both showed antigen binding when combined with cL or kgL221A, the kgH341A gene with all 11 changes appearing to be the superior choice.

15 3 INTERIM CONCLUSIONS

It has been demonstrated, therefore, for OKT3 that to transfer antigen binding ability to the humanised antibody, mouse residues outside the CDR regions defined by the Kabat hypervariability or structural loop choices are required for both the light and heavy chains. Fewer extra residues are needed for the

- light chain, possibly due to the higher initial homology between the mouse and human kappa variable regions.
- Of the changes seven (1 and 3 from the light chain and 6, 23, 71, 73 and 76 from the heavy chain) are predicted from a knowledge of other antibody structures to be either partly exposed or on the antibody surface. It has been shown here that residues 1 and 3 in the light chain are not absolutely required to be the mouse sequence; and for the heavy chair the grt3418 heavy chain in combination with the 2214 light chain generated only weak binding activity. Therefore the presence of the 6, 23 and 24 changes are important to maintain a binding affinity similar to that of the murine autitody; at was important, therefore, be further study the individual contribution of othe other 8 mouse residues of
- the kgH341A gene compared to kgH341.

  16. FURTHER CDR-GRAFTING EXPERIMENTS

- Additional CDR-grafted heavy chain genes were prepared substantially as described above. With reference to Table 2 the further heavy chain genes were based upon the gh341 (plasmid pJA178) and gh341A (plasmid pJA185) with either mouse OKT3 or human KOL residues at 6, 22, 24, 48, 49, 63, 71, 73, 76, 78, 88 and 91, as indicated. The CDR-grafted light chain genes used in these further experiments
- 73, 76, 78, 88 and 91, as indicated. The CDR-grafted light chain genes used were gL221, gL221A, gL221B and gL221C as described above.

## TABLE 2

## OKT3 HEAVY CHAIN CDR GRAFTS

## 1. gH341 and derivatives

10	RES NUM	6	23	24	48	49	63	71	73	76	78	88	91
	OKT3vh	9	ĸ	Α.	1	G	F	Ţ	K	S	Α.	A	<u> </u>
	gH341	E	s	s	v	A	F	R	N	N	L	G	F JA178
	gH341A	9	K	A	I	G	v	T	K	S	٨	A	Y JA185
15	gH341E	9	к	Α_	_1	G	v	<u>T</u>	ĸ	S	^	G	G JA198
	gH341*	0	K	Α.	1	G	٧	<u>T</u>	K	N	Δ	G	F JA207
	gH341*	9	K	Α.	1	G	v	R	ĸ	N	Α	C	F JA209
20	gH341D	<u>Q_</u>	К	· A	1	G	v	Ţ_	K	N	L	G	F JA197
	gH341*	0	K	A	1	G	v	R	N	N	L	G	F JA199
	gH341C	9	K		v	A	Ē.	R	N	N	L	G	F JA184
	gH341*	Q	5	A	1	G	٧	Ţ	K	5	A		Y JA203
25	gH341*	E	s	<u>A</u>	1	G	v	Ţ	K		٨	Α.	Y JA205
	gH341B	E	s	s	1	G	v	T	K	5	_^	Α	<u>Y</u> JA183
	gH341*	Q	s	<u>A</u>		G	v	<u>T</u>	K	5		G	F JA204
	gH341*	E	s	Α	1	G	v	Ţ	ĸ	S	A	G	F JA206
30	gH341*	Q	S	A	1	Ç	v	<u>T</u>	<u> </u>	N	Ž.	G	F JA208
	KOL	E	s	s	v	Α		R	N	N	L	G	F

## 35 OKT3 LIGHT CHAIN CDR GRAFTS

## 2. gL221 and derivatives

40	RES NUM	1	3	46	47
	OKT3v1	Q	٧	R	U
	GL221	D	Q	L	L DA221
45	gL221A	٩	ν	R	W DA221
	gL221B	9_	v	L	L DA221
	GL221C	D	Q	R	₩ DA221
	RE1	D	0	1.	L

# MURINE RESIDUES ARE UNDERLINED

The CDR-grafted heavy and light chain genes were co-expressed in COS cells either with one another in various combinations but also with the corresponding murine and chimeric heavy and light chain genes substantially as described above. The resultant antibody products were then assayed in binding and blocking assays with HPB-ALL cells as described above.

The results of the assays for various grafted heavy chains co-expressed with the gL221C light chain are given in Figures 7 and 8 (for the JA184, JA185, JA197 and JA198 constructs - see Table 2), in Figure 9 (for the JA183, JA184, JA185 and JA197 constructs) in Figure 10 (for the chimeric, JA185, JA198, JA204, JA205, JA207, JA208 and JA209 constructs) and in Figure 11 (for the JA183, JA184, JA185, JA198, JA203, SLA905, JA207, JA208 and JA208 constructs).

The basic grafted product without any human to murine changes in the variable frameworks, i.e. gl.221 co-expressed with gh341 (JA178), and also the "fully grafted" product, having most human to murine changes in the grafted heavy chain framework, i.e. gl.221C co-expressed with gh341A (JA185), were assayed for relative binding affinity in a competition assay against murine OKT3 reterence standard, using 14PB-ALL colls. The assay used was as described above in section 3.3. The results obtained are given in Figure 12 for the basic grafted product has neglibble binding ability as compared with the OKT3 murine reference standard; whereas the "fully grafted" product has a binding ability very similar to that of the OKT3 murine reference standard.

15 The binding and blocking assay results indicate the following:

The JA196 and JA207 constructs appear to have the best binding characteristics and similar binding abilities, both substantially the same as the chimeric and fully grafted gH341A products. This indicates that positions 88 and 91 and position 76 are not highly critical for maintaining the OKT3 binding ability; whereas at least some of positions 6, 23, 24, 48, 49, 71, 73 and 78 are more important.

This is borne out by the finding that the JA209 and JA199, although of similar binding ability to one another, are of lower binding ability than the JA198 and JA207 constructs. This indicates the importance of having mouse residues at positions 71, 73 and 78, which are either completely or partially human in the JA199 and JA209 constructs respectively.

Moreover, on comparing the results obtained for the JA205 and JA183 constructs it is seen that there is a decrease in binding going from the JA205 to the JA183 constructs. This indicates the importance of rotating a mouse residue at position 23, the only position changed between JA205 and JA183.

These and other results lead us to the conclusion that of the 11 mouse framework residues used in the gH341A (ARI85) construct, it is important to retain mouse residues at all of positions 6, 23, 24, 48 and 49, and possibly for maximum binding affinity at 71, 73 and 78.

Similar Experiments were carried out to CDR-graft a number of the rodent antibodies including antibodies having specificity for CD4 (OKT4), ICAM-1 (R6-5), TAG72 (B72.3), and TNFa(61E71, 101.4, hTNF1, hTNF2 and hTNF3).

#### **EXAMPLE 2**

# CDR-GRAFTING OF A MURINE ANTI-CD4 T CELL RECEPTOR ANTIBODY, OKT4A

#### 45 THE LIGHT CHAIN

The human acceptor framework used for the grafted light chains was RE1. The preferred LCDR2 light chain has human to mouse changes at positions 33, 34, 38, 49 and 99 in addition to the structural loop CDRs. Of these changed positions, positions 33, 34 and 89 fall within the preferred extended CDRs of the present invention (positions 33 and 34 in CDR1 and position 89 in CDR3).

The human to murine changes at positions 38 and 49 corresponds to positions at which the amino acid residues are preferably donor murine amino acid residues in accordance with the present invention.

A comparison of the amino acid sequences of the donor murine light chain variable domain and the RET munan acceptor light chain variable further reveals that the murine and human residues are identical at all 50 of positions 46, 48 and 71 and at all of positions 2, 4, 6, 35, 36, 44, 47, 62, 64-68, 85, 87, 99, 99 and 101 and 102. However the amino acid residue at position 58 in LCDR2 is the human RET framework residue not the mouse OKT4 residue as would be preferred in accordance with the present invention.

#### THE HEAVY CHAIN

The human acceptor framework used for the grafted heavy chains was KOL.

The preferred CDR graft HCDR10 heavy chain has human to mouse changes at positions 24, 35, 57, 58, 50, 88 and 91 in addition to the structural loop CDRs.

Of these positions, positions 35 (CDR1) and positions 57, 58 and 69 (CDR2) fall within the preferred extended CDRs of the present invention. Also the human to mouse change at position 24 corresponds to a position at which the amino acid residue is a donor murine residue in accordance with the present invention. Moreover, the human to mouse changes at positions 88 and 91 correspond to positions at which the amino acid residues are optionally donor murine residues.

Moreover, a comparison of the murino OKT4A and human KOL heavy chain variable amino acid sequences reveals that the murine and human residues are identical at all of positions 23, 49, 71, 73 and 78 and at all of positions 2, 4, 6, 25, 68, 73, 94, 74, 89, 94, 103, 104, 106 and 107.

Thus the OKT4A CDR-grafted heavy chain HCDR10 corresponds to a particularly preferred embodiment according to the present invention.

#### **EXAMPLE 3**

## CDR-GRAFTING OF AN ANTI-MUCIN SPECIFIC MURINE ANTIBODY, B72.3

The cloning of the genes coding for the anti-mucin specific murine monoclonal antibody B72.3 and the preparation of B72.3 mouse-human chimeric antibodies has been described previously (ref. 13 and WO 8901783). CDR-grafted versions of B72.3 were propared as follows.

#### (a) B72.3 Light Chain

CDR-grafting of this light chain was accomplished by direct transfer of the murine CDRs into the framework of the human light chain RE1. The regions transferred were:

	CDR Number	Residues
Į	1	24-34
	2	50-56
	9	90-98

The activity of the resulting grafted light chain was assessed by co-expression in COS cells, of genes for the combinations: B723 ethB72.3 ct.

## and B72.3 cH/B72.3 gL

Supernatants were assayed for antibody concentration and for the ability to bind to microtitre plates of coated with mucin. The results obtained indicated that, in combination with the B72.3 c.H chain, B72.3 c.L and B72.3 c.H as similar binding properties.

Comparison of the murine B72.3 and REI light chain amino acid sequences reveals that the residues are identical at positions 46, 58 and 71 but are different at position 48. Thus changing the human residue to the donor mouse residue at position 48 may further improve the binding characteristics of the CDR-grafted Light chain, (B72.3 gL) in accordance with the present invention.

## (b) B72.3 heavy chain

## i. Choice of framework

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At the outset it was necessary to make a choice of human framework. Simply put, the question was as follows: Was it necessary to use the framework regions from an antibody whose crystal structure was known or could be choice be made on some other critical?

For B7.23 heavy chain, it was reasoned that, while knowledge of structure was important, transfer of the CDRs from mouse to human frameworks might be tacilitated if the overall homology between the donor and receptor trameworks was maximised. Comparison of the B7.23 heavy chain sequence with tiose in Kabat (ref. 4) for human heavy chains showed clearly that B7.23 heavy chain sequence with and NEVMI (for which crystal structures are available) but was very homologous to the heavy chain

On this basis, EU was chosen for the CDR-grafting and the following residues transferred as CDRs.

CDR Number	Residues
1	27-36
2	50-63
3	93-102

Also it was noticed that the FR4 region of EU was unlike that of any other human (or mouse) antibody. Consequently, in the grafted heavy chain genes this was also changed to produce a "consensus" human sequence. (Preliminary experiments showed that grafted heavy chain genes containing the EU FR4 sequence expressed very poorly in transient expression systems.)

## ii. Results with grafted heavy chain genes

Expression of grafted heavy chain genes containing all human framework regions with either gL or cL genes produced a grafted antibody with little ability to bind to mucin. The grafted antibody had about 1% the activity of the chimeric antibody, in these experiments, however, it was noted that the activity of the grafted antibody could be increased to - 10% of 872.3 by exposure to pits of 2-3.5.

to the granted amount of the control of the granted antibody could be improved without acid treatment, it was possulated that acid exposure brought about the protonation of an acidic residue (pfx of aspartic acid = 3.86 and of glutamine acid = 4.25) which in turn caused a change in structure of the CDR loops, or allowed better access of antigen.

From comparison of the sequences of B723 (ref. 13) and EU (refs. 4 and 5), it was clear that, in going from the mouse to human frameworks, only two positions had been changed in such a way that acidic residues had been introduced. These positions are at residues 73 and 81, where K to E and Q to E changes had been made, respectively.

Which of these positions might be important was determined by examining the crystal structure of the KOL antibody. In KOL heavy chain, position 81 is far removed from either of the CDR loops.

Position 73, however, is close to both CDRs 1 and 3 of the heavy chain and, in this position it was possible to envisage that a K to E change in this region could have a detrimental effect on antigen binding.

#### iii. Framework changes in B72.3 gH gene

On the basis of the above analysis, £73 was mutated to a lysine (K), It was found that this change had a dramatic effect on the ability of the grafted Ab to bind to mucin. Further the ability of the grafted 8723 produced by the mutated gHgL combination to bind to mucin was similar to that of the B723 chimaric antibody.

### iv. Other framework changes

In the course of the above experiments, other changes were made in the heavy chain framework regions. Within the accuracy of the assays used, none of the changes, either alone or together, appeared beneficial.

## v. Other

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All assays used measured the ability of the grafted Ab to bind to mucin and, as a whole, indicated that the single framework change at position 73 is sufficient to generate an antibody with similar binding properties to 872.9.

Comparison of the B72.3 murine and EU heavy chain sequences reveals that the mouse and human residues are identical at positions 23, 24, 71 and 78.

Thus the mutated CDR-grafted B72.3 heavy chain corresponds to a preferred embodiment of the present invention.

#### EXAMPLE 4

## CDR-GRAFTING OF A MURINE ANTI-ICAM-1 MONOCLONAL ANTIBODY

A murine antibody, R6-5-D6 (EP 0314863) having specificity for Intercellular Adhesion Molecule 1 (ICAM-1) was CDR-grafted substantially as described above in previous examples. This work is described in greater detail in co-pending application, British Patent Application No. 9009549.8, the disclosure of which is incorporated herein by reference.

The human EU framework was used as the acceptor framework for both heavy and light chains. The CDRgrafted antibody currently of choice is provided by co-expression of grafted light chain gL221A and grafted heavy chain gH341D which has a binding affinity for ICAM 1 of about 75% of that of the corresponding

mouse-human chimeric antibody.

## LIGHT CHAIN

gL221A has murine CDRs at positions 24-34 (CDR1), 50-56 (CDR2) and 89-97 (CDR3). In addition several framework residues are also the murine amino acid. These residues were chosen after consideration of the possible contribution of these residues to domain packing and stability of the conformation of the artitigen binding region. The residues which have been retained as mouse are at positions 2, 3, 48 (?), 60, 84, 85 and 87. Comparison of the murine anti-LRM1 and human EU light chain amino acid sequences reveals that the murine and human residues are identical at positions 46, 58 and 71.

### HEAVY CHAIN

pt341D has murine CDRs at positions 28-53 (CDR1), 50-56 (CDR2) and34-1008 (CDR3). In addition murine residuous were used in ght41D at positions 24, 48, 69, 71, 73, 80, 88 and 91. Comparison of the murine arti-ICAM 1 and human EU heavy chain amino acid sequences are identical at positions 23, 49 and 78.

#### EXAMPLE 5

#### .....

## CDR-Grafting of murine anti-TNF2 antibodies

A number of murine anti-TNFa monoclonal antibodies were CDR-grafted substantialty as described above in previous examples. These antibodies include the murine monoclonal antibodies designated 61 EF1, hTNF1, hTNF3 and 101.4 A brief summary of the CDR-grafting of each of these antibodies is given below.

## 30 61E71

A similar analysis as described above (Example 1, Section 12.1) was done for 81E71 and for the heavy chain 10 residues were identified at 23, 24, 48, 48, 68, 69, 71, 73, 75 and 88 as residues to potentially retain as murine. The human frameworks chosen for CDR-grefting of this antibody, and the hTNR3 and 31 01.4 antibodies were RE1 for the fight chain and KOL for the heavy chain.

Three genes were built, the first of which contained 23, 24, 48, 49, 71 and 73 (gli341(6)) as murine residues. The second gene also had 75 and 88 as murine residues (gli341(10)) while the third gene additionally had 68, 69, 75 and 88 as murine residues (gli341(10)). Each was co-expressed with gl221 hgli341(6) and gl221(gli341(4)) and gl221(gli341(4)) and gl221(gli341(4)) and gl221(gli341(4)) and gl231(gli341(6)) and gl321(gli341(6)) and gl3

40 as well to TNF as murine 61E71. The gt.221/gH341(10) antibody did not express and this combination was not taken further. Subsequently the gt.221/gH341(6) antibody was assessed in an L929 cell competition assay in which the antibody competes against the TNF receptor on 1292 cells for binding to TNF in solution. In this assay the gt.221(gH341(6) antibody was approximately 10% as active as murine 61E71.

### hTNF1

hTNF1 is a monoclonal antibody which recognises an epitope on human TNF- . The EU human framework was used for CDR-grafting of both the heavy and light variable domains.

## Heavy Chain

In the CDR-grafted heavy chain (ghTNF1) mouse CDRs were used at positions 26-35 (CDR1), 50-65 (5DR2) and 95-102 (CDR3), Mouse residues were also used in the frameworks at positions 48, 67, 68, 71, 73, 76, 89, 91, 94 and 108. Comparison of the TNF1 mouse and EU human heavy chain residues reveals that these are identical at ostitions 23, 24, 29 and 78.

## Light Chain

In the CDR-grafted light chain (gLhTNF1) mouse CDRs wre used at positions 24-34 (CDR1), 50-56 (CDR2) and 89-97 (CDR3). In addition mouse residues were used in the frameworks at positions 3, 42, 48, 5 49, 83, 106 and 108. Comparison of the hTNF1 mouse and EU human light chain residues reveals that these are identical at positions 46, 58 and 71.

The grafted hTNF1 heavy chain was co-expressed with the chimeric light chain and the binding ability of the product compared with that of the chimeric light chain/chimeric heavy chain product in a TNF binding assay. The grafted heavy chain product appeared to have binding ability for TNF slightly better than the 10 fully chimeric product.

Similarly, a grafted heavy chain/grafted light chain product was co-expressed and compared with the fully chimeric product and found to have closely similar binding properties to the latter product.

## hTNF3

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hTNF3 recognises an epitope on human TNF-a. The sequence of hTNF3 shows only 21 differences compared to 61E71 in the light and heavy chain variable regions, 10 in the light chain (2 in the CDRs at positions 50, 96 and 8 in the framework at 1, 19, 40, 45, 46, 76, 103 and 106) and 11 in the heavy chain (3 in the CDR regions at positions 52, 60 and 95 and 8 in the framework at 1, 10, 38, 40, 67, 73, 87 and 105). 20 The light and heavy chains of the 61E71 and hTNF3 chimeric antibodies can be exchanged without loss of activity in the direct binding assay. However 61E71 is an order of magnitude less able to compete with the TNF receptor on L929 cello for TNF-a compared to hTNF3. Based on the 61E71 CDR grafting data gL221 and gH341(+23, 24, 48, 49 71 and 73 as mouse) genes have been built for hTNF3 and tested and the resultant grafted antibody binds well to TNF-a, but competes very poorly in the L929 assay. It is possible 25 that in this case also the framework residues identified for OKT3 programme may improve the competitive binding ability of this antibody.

## 101.4

101.4 is a further murine monoclonal antibody able to recognise human TNF-a. The heavy chain of this antibody shows good homology to KOL and so the CDR-grafting has been based on RE1 for the light chain and KOL for the heavy chain. Several grafted heavy chain genes have been constructed with conservative choices for the CDR's (gH341) and which have one or a small number of non-CDR residues at positions 73, 78 or 77-79 inclusive, as the mouse amino acids. These have been co-expressed with cL or gL221. In all as cases binding to TNF equivalent to the chimeric antibody is seen and when co-expressed with cL the resultant antibodies are able to compete well in the L929 assay. However, with gL221 the resultant antibodies are at least an order of magnitude less able to compete for TNF against the TNF receptor on L929 cells.

Mouse residues at other positions in the heavy chain, for example, at 23 and 24 together or at 76 have 40 been demonstrated to provide no improvement to the competitive ability of the grafted antibody in the L929 assay.

A number of other antibodies including antibodies having specificity for interleukins e.g. IL1 and cancer markers such as carcinoembryonic antigen (CEA) e.g. the monoclonal antibody A5B7 (ref. 21), have been successfully CDR-grafted according to the present invention.

45 It will be appreciated that the foregoing examples are given by way of illustration only and are not intended to limit the scope of the claimed invention. Changes and modifications may be made to the methods described whilst still falling within the spirit and scope of the invention.

### References

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## SEQUENCE LISTING

	•
	(1) GENERAL INFORMATION:
5	(i) APPLICANT: (A) NAME: CELLTECH LIMITED
	(B) STREET: 216 BATH ROAD (C) CITY: SLOUGH (D) STATE: BERKSHIRE
10	(E) COUNTRY: UNITED KINGDOM (P) POSTAL CODE (ZIP): SL1 4EN (G) TELEPHONE: 0753 534655
	(H) TELEFAX: 0753 536632 (I) TELEX: 848473
	(ii) TITLE OF INVENTION: HUMANISED ANTIBODIES
15	(iii) NUMBER OF SEQUENCES: 33
	(iv) COMPUTER READALE FORM:  (A) MEDIUM TYPE: Floppy disk  (B) COMPUTER: IBM PC compatible  (C) OPERATING SYSTEM: PC-DOS/MS-DOS  (D) SOFTMARE: PatentIn Release #1.0, Version #1.25
20	(EPO)
	(2) INFORMATION FOR SEQ ID NO: 1:
25	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANGENESS: single (D) TOPOLOSY: linear
30	(ii) MOLECULE TYPE: cDNA
-	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:
	TCCAGATGTT AACTGCTCAC
35	(2) INFORMATION FOR SEQ ID NO: 2:
	(i) SEQUENCE CHARACTERISTICS:
40	(A) LENGTH: 23 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
	(ii) MOLECULE TYPE: cDNA
45	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:
	CAGGGGCCAG TGGATGGATA GAC 23
	(2) INFORMATION FOR SEQ ID NO: 3:
50	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 9 amino acids (B) TYPE: amino acid

(C) STRANDEDNESS: single (D) TOPOLOGY: linear

	(ii) MOLECULE TYPE: protein -
5	(v) FRAGMENT TYPE: internal
	(xi) SEQUENCE DESCRIPTION: SEQ ID No: 3:
10	Leu Glu Ile Asn Arg Thr Val Ala Ala 1
	(2) INFORMATION FOR SEQ ID NO: 4:
15	(i) SEQUENCE CHARACTERISTICS: (A) LENCTH: 943 base pairs (B) TFFE: mucleic acid (C) STRANDENNESS: single (D) TOPOLOGY: linear
	(ii) MOLECULE TYPE: cDNA
20	
	(x1) SEQUENCE DESCRIPTION: SEQ ID NO: 4:
	GAATTCCCAA AGACAAAATG GATTTTCAAG TGCAGATTTT CAGCTTCCTG CTAATCAGTG 60
25	CCTCAGTCAT AATATCCAGA GGACAAATTG TTCTCACCCA GTCTCCAGCA ATCATGTCTG 120
	CATCTCCAGG GGAGAAGGTC ACCATGACCT GCAGTGCCAG CTCAAGTGTA AGTTACATGA 180
30	ACTGGTACCA GCAGAAGTCA GGCACCTCCC CCAAAAGATG GATTTATGAC ACATCCAAAC 240
	TGGCTTCTGG AGTCCCTGCT CACTTCAGGG GCAGTGGGTC TGGGACCTCT TACTCTCTCA 300
35	CAATCAGCGG CATGGAGGCT GAAGATGCTG CCACTTATTA CTGCCAGCAG TGGAGTAGTA 360
	ACCCATTCAC GTTCGGCTCG GGGACAAAGT TGGAAATAAA CCGGGCTGAT ACTGCACCAA 420
40	CTGTATCCAT CTTCCCACCA TCCAGTGAGC AGTTAACATC TGGAGGTGCC TCAGTCGTGT 480
	GCTTCTTGAA CAACTTCTAC CCCAAAGACA TCAATGTCAA GTGGAAGATT GATGGCAGTG 540
45	AACGACAAAA TGGCGTCCTG AACAGTTGGA CTGATCAGGA CAGCAAAGAC AGCACCTACA 600
	GCATGAGCAG CACCCTCACG TTGACCAAGG ACGAGTATGA ACGACATAAC AGCTATACCT 660
50	GTGAGGCCAC TCACAAGACA TCAACTTCAC CCATTGTCAA GAGCTTCAAC AGGAATGAGT 720
-	GTTAGAGACA AAGGTCCTGA GACGCCACCA CCAGCTCCCA GCTCCATCCT ATCTTCCCTT 780
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CTAAGGTC	TT G	GAGG	CTTC	c cc	ACAA	GCGC	TTA	CCAC	rgt :	PGCG	TGC:	rc T	AAAC	CTCC	r	84
CCCACCTC	CT T	CTCC	rccr(	с ст	CCCT	PTCC	TTG	CTT	TA :	PCAP(	CTA	À A	TTTG:	caga	A	90
AATATTCA	AT A	AAGT	GAGT	TT	rgcc	<b>TTGA</b>	AAA	AAAA	AA.	AAA						94
	SEQUAL (A)	JENC: LEI TYI STI	E CHANGTH	ARAC 23: amin EDNE:	reri 3 am 5 ac SS:	STIC ino id sing ar	S: acid:	5								
	SEQ						no Ti	. 110						٠.		
	yab									Île	Ser	Ala	Ser	Val 15	Ile	
Ile	Ser	Arg	Gly Gly	Gln	Ile	Val	Leu	Thr 25	Gln	Ser	Pro	Ala	Ile 30	Met	Ser	
λla	Ser	Pro 35	Gly	Glu	Lys	Val	Thr 40	Met	Thr	Cys	Ser	Ala 45	Ser	Ser	Ser	
	Ser 50					55					60					
65	Trp				70					75			•	,	80	
	Glu			85					90					95		
	Pro		100					105					110			
	Thr	115					120					125				
Ī	130					135					140					
145	Ser				150					155					160	
-	Asp			165		-			170	-			Ī	175		
	Val		180					185					190			
	Met	195					200					205				
	Ser 210 Lys	•		-		215			rys	rnr	Ser 220	rnr	ser	rro	TTG	
225					230			CYS								
(2) INFO	RMAT:	EON I	FOR S	SEQ :	ID NO	): 6	:									

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 1570 base pairs (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

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## (ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6: 10 GRATTCCCCT CTCCACAGAC ACTGAAAACT CTGACTCAAC ATGGAAAGGC ACTGGATCTT 60 TCTACTCCTG TTGTCAGTAA CTGCAGGTGT CCACTCCCAG GTCCAGCTGC AGCAGTCTGG 120 GGCTGAACTG GCAAGACCTG GGGCCTCAGT GAAGATGTCC TGCAAGGCTT CTGGCTACAC 180 CTTTACTAGG TACACGATGC ACTGGGTAAA ACAGAGGCCT GGACAGGGTC TGGAATGGAT 240 TGGATACATT AATCCTAGCC GTGGTTATAC TAATTACAAT CAGAAGTTCA AGGACAAGGC 300 CACATTGACT ACAGACAAAT CCTCCAGCAC AGCCTACATG CAACTGAGCA GCCTGACATC 360 TGAGGACTET GCAGTETATT ACTGTGCAAG ATATTATGAT GATCATTACT GCCTTGACTA 420 CTGGGGCCAA GGCACCACTC TCACAGTCTC CTCAGCCAAA ACAACAGCCC CATCGGTCTA 480 TCCACTGGCC CCTGTGTGTG GAGATACAAC TGGCTCCTCG GTGACTCTAG GATGCCTGGT 540 CAAGGGTTAT TTCCCTGAGC CAGTGACCTT GACCTGGAAC TCTGGATCCC TGTCCAGTGG 600 TGTGCACACC TTCCCAGCTG TCCTGCAGTC TGACCTCTAC ACCCTCAGCA GCTCAGTGAC 660 TGTAACCTCG AGCACCTGGC CCAGCCAGTC CATCACCTGC AATGTGGCCC ACCCGGCAAG 720 CAGCACCAAG GTGGACAAGA AAATTGAGCC CAGAGGGCCC ACAATCAAGC CCTGTCCTCC 780 30 ATGCAAATGC CCAGCACCTA ACCTCTTGGG TGGACCATCC GTCTTCATCT TCCCTCCAAA 840 GATCAAGGAT GTACTCATGA TCTCCCTGAG CCCCATAGTC ACATGTGTGG TGGTGGATGT 900 GAGCGAGGAT GACCCAGATG TCCAGATCAG CTGGTTTGTG AACAACGTGG AAGTACACAC 960 AGCTCAGACA CAAACCCATA GAGAGGATTA CAACAGTACT CTCCGGGTGG TCAGTGCCCT 1020 35 CCCCATCCAG CACCAGGACT GGATGAGTCC CAAGGAGTTC AAATGCAAGG TCAACAACAA 1080 AGACCTCCCA CCCCCATCG AGAGACCAT CTCAAAACCC AAAGGGTCAG TAAGAGCTCC 1140 ACAGGRAPAT GTCTTGCCTC CACCAGAAGA AGAGATGACT AAGAAACAGG TCACTCTGAC 1200 40 CTGCATGGTC ACAGACTTCA TGCCTGAAGA CATTTACGTG GAGTGGACCA ACAACGGGAA 1260 AACAGAGCTA AACTACAAGA ACACTGAACC AGTCCTGGAC TCTGATGGTT CTTACTTCAT 1320 GTACAGCAAG CTGAGAGTGG AAAAGAAGAA CTGGGTGGAA AGAAATAGCT ACTCCTGTTC 1380 45 AGTGGTCCAC GAGGGTCTGC ACAATCACCA CACGACTAAG AGCTTCTCCC GGACTCCGGG 1440 TARATGAGCT CAGCACCCAC ARAACTCTCA GGTCCAAAGA GACACCCACA CTCATCTCCA 1500 1560

(2) INFORMATION FOR SEQ ID NO: 7:

AAAGGAATTC

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- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 468 amino acids
    (B) TYPE: amino acid
    (C) STRANDEDNESS: single
  - (C) STRANDEDNESS: sing (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7: 10 Met Glu Arg His Trp Ile Phe Leu Leu Leu Leu Ser Val Thr Ala Gly Val His Ser Gln Val Gln Leu Gln Gln Ser Gly Ala Glu Leu Ala Arg 20 25 30 Pro Gly Ala Ser Val Lys Met Ser Cys Lys Ala Ser Gly Tyr Thr Phe  $35 \hspace{1cm} 40 \hspace{1cm} 45$ Thr Arg Tyr Thr Met His Trp Val Lys Gln Arg Pro Gly Gln Gly Leu 50 60 Glu Trp Ile Gly Tyr Ile Asn Pro Ser Arg Gly Tyr Thr Asn Tyr Asn 65 75 80 Gln Lys Phe Lys Asp Lys Ala Thr Leu Thr Thr Asp Lys Ser Ser Ser 85 90 95 Thr Ala Tyr Met Gln Leu Ser Ser Leu Thr Ser Glu Asp Ser Ala Val Tyr Tyr Cys Ala Arg Tyr Tyr Asp Asp His Tyr Cys Leu Asp Tyr Trp Gly Gln Gly Thr Thr Leu Thr Val Ser Ser Ala Lys Thr Thr Ala Pro 130 135 140 Ser Val Tyr Pro Leu Ala Pro Val Cys Gly Asp Thr Thr Gly Ser Ser 145 150 155 160 Val Thr Leu Gly Cys Leu Val Lys Gly Tyr Phe Pro Glu Pro Val Thr Leu Thr Trp Asn Ser Gly Ser Leu Ser Ser Gly Val His Thr Phe Pro Ala Val Leu Gln Ser Asp Leu Tyr Thr Leu Ser Ser Ser Val Thr Val Thr Ser Ser Thr Trp Pro Ser Gln Ser Ile Thr Cys Asn Val Ala His 210 220 Pro Ala Ser Ser Thr Lys Val Asp Lys Lys Ile Glu Pro Arg Gly Pro 225 230 235 240 Thr Ile Lys Pro Cys Pro Pro Cys Lys Cys Pro Ala Pro Asn Leu Leu 245 250 255 Gly Gly Pro Ser Val Phe Ile Phe Pro Pro Lys Ile Lys Asp Val Leu 260 265 270 Met Ile Ser Leu Ser Pro Ile Val Thr Cys Val Val Val Asp Val Ser 275 280 285

Glu Asp Asp Pro Asp Val Gln Ile Ser Trp Phe Val Asn Asn Val Glu

			290					295					300				
		Val 305	His	Thr	Ala	G1n	Thr 310	Gln	Thr	His	Arg	61u 315	Asp:	љуr	Asn	Ser	Thr 320
		Leu	Arg	Val	Val	Ser 325	Ala	Leu	Pro	Ile	Gln 330	His	Gln	Asp	Trp	Met 335	Ser
		Gly	Lys	Glu	Phe 340	Lys	Cys	Lys	Va1	Asn 345	Asn	Lys	Asp	Leu	Pro 350	Ala	Pro
		Ile	Glu	Arg 355	Thr	Ile	Ser	Lys	Pro 360	Lys	Gly	Ser	Val	Arg 365	Ala	Pro	G1n
		Val	Tyr 370	Va1	Leu	Pro	Pro	Pro 375		Glu	Glu	Met	Thr 380	Lys	Lys	Gln	Val
;		Thr 385	Leu	Thr	Cys	Met	Val 390	Thr	Asp	Phe	Met	Pro 395	Glu	Asp	Ile	Tyr	Val 400
		Glu	Trp	Thr	Asn	Asn 405	Gly	Lys	Thr	Glu	Leu 410	Asn	Tyr	Lys	Asn	Thr 415	Glu
,		Pro	Val	Leu	Asp 420	ser.	Asp	Gly	Ser	Tyr 425	Phe	Met	Tyr	Ser	Lys 430	Leu	Arg
		Val	Glu	<b>Lys</b> 435	Lys	Asn	Trp	Val	Glu 440	Arg	Asn	Ser	Tyr	Ser 445	Cys	Ser	Val
<b>.</b>		Val	His 450	Glu	Gly	Leu	His	Asn 455	His	His	Thr	Thr	Lys 460	Ser	Phe	Ser	Arg
		Thr 465	Pro	Gly	Lys		٠										
	(2)	INPOF	(TAM	ON E	or s	EQ 1	D NO	: 8:							-		
9		(i)	(A) (B) (C)	LENCE TYI STF	GTH: PE: a VANDE	107 mino DNES	ami aci s: s	no a d ingl	cids	:							
5		(ii)	MOLE	CULE	TYE	E: I	rote	in									

Phe Gly Ser Gly Thr Lys Leu Glu Ile Asn Arg

- (2) INFORMATION FOR SEQ ID NO: 9:
  - MICHIELDA FOR SEG ID NO. 3.
  - (i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 108 amino acids
    - (B) TYPE: amino acid (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein

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- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:
- Asp Ile Gln Met Thr Gln Ser Pro Ser Ser Leu Ser Ala Ser Val Gly 1 10 15
- Asp Arg Val Thr Ile Thr Cys Gln Ala Ser Gln Asp Ile Ile Lys Tyr 20 25 30
- Leu Asn Trp Tyr Gln Gln Thr Pro Gly Lys Ala Pro Lys Leu Leu Ile  $35 \hspace{1.5cm} 40 \hspace{1.5cm} 45$
- Tyr Glu Ala Ser Asn Leu Gln Ala Gly Val Pro Ser Arg Phe Ser Gly 50 60
- Ser Gly Ser Gly Thr Asp Tyr Thr Phe Thr Ile Ser Ser Leu Gln Pro  $^{65}$
- Glu Asp Ile Ala Thr Tyr Cys Gln Gln Tyr Gln Ser Leu Pro Tyr 85 90 95
- Thr Phe Gly Gln Gly Thr Lys Leu Gln Ile Thr Arg
- (2) INFORMATION FOR SEQ ID NO: 10:
  - (i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 119 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10:
  - Gln Val Gln Leu Gln Gln Ser Gly Ala Glu Leu Ala Arg Pro Gly Ala 1 10 15
  - Ser Val Lys Met Ser Cys Lys Ala Ser Gly Tyr Thr Phe Thr Arg Tyr 20 25 30
  - Thr Met His Trp Val Lys Gln Arg Pro Gly Gln Gly Leu Glu Trp Ile 35 40 45
- Gly Tyr Ile Asn Pro Ser Arg Gly Tyr Thr Asn Thr Asn Gln Lys Phe
- Lys Asp Lys Ala Thr Leu Thr Thr Asp Lys Ser Ser Ser Thr Ala Tyr 65 70 75 80

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Met Gln Leu Ser Ser Leu Thr Ser Glu Asp Ser Ala Val Tyr Tyr Cys Ala Arg Tyr Tyr Asp Asp His Tyr Cys Leu Asp Tyr Trp Gly Gln Gly 100 105 110 Thr Thr Leu Thr Val Ser Ser (2) INFORMATION FOR SEQ ID NO: 11: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 126 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: protein 15 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 11: Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 10 15 20 Ser Leu Arg Leu Ser Cys Ser Ser Ser Gly Phe Ile Phe Ser Ser Tyr Ala Met Tyr Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 25 Ala Ile Ile Trp Asp Asp Gly Ser Asp Gln His Tyr Ala Asp Ser Val Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Phe 65 70 75 80 30 Leu Gln Met  $\lambda$ sp Ser Leu  $\lambda$ rg Pro Glu  $\lambda$ sp Thr Gly Val Tyr Phe Cys 85 90 95 Ala Arg Asp Gly Gly His Gly Phe Cys Ser Ser Ala Ser Cys Phe Gly 100 105 110Pro Asp Tyr Trp Gly Gln Gly Thr Pro Val Thr Val Ser Ser (2) INFORMATION FOR SEQ ID NO: 12: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 119 amino acids (B) TYPE: amino acid 40 (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: protein (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12: Gln Val Gln Leu Gln Gln Ser Gly Ala Glu Leu Ala Arg Pro Gly Ala Ser Val Lys Met Ser Cys Lys Ala Ser Gly Tyr Thr Phe Thr Arg Tyr 55

Thr Met His Trp Val Lys Gln Arg Pro Gly Gln Gly Leu Glu Trp Ile Gly Tyr Ile Asn Pro Ser Arg Gly Tyr Thr Asn Tyr Asn Glm Lys Phe Lys Asp Lys Ala Thr Leu Thr Thr Asp Lys Ser Ser Ser Thr Ala Tyr Met Gln Leu Ser Ser Leu Thr Ser Glu Asp Ser Ala Val Tyr Tyr Cys 10 Ala Arg Tyr Tyr Asp Asp His Tyr Cys Leu Asp Tyr Trp Gly Gln Gly 100 105 110 Thr Thr Leu Thr Val Ser Ser 15 (2) INFORMATION FOR SEQ ID NO: 13: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 119 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single 20 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: protein (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 13: Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 10 15 Ser Leu Arg Leu Ser Cys Ser Ser Ser Gly Tyr Thr Phe Thr Arg Tyr Thr Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val Ala Tyr Ile Asn Pro Ser Arg Gly Tyr Thr Asn Tyr Asn Gln Lys Phe Lys Asp Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr Leu Phe 65 70 75 80

(2) INFORMATION FOR SEQ ID NO: 14:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 119 amino acids

(B) TYPE: amino acid (C) STRANDEDNESS: single

(D) TOPOLOGY: linear

Thr Thr Leu Thr Val Ser Ser

(ii) MOLECULE TYPE: protein

Leu Gln Met Asp Ser Leu Arg Pro Glu Asp Thr Gly Val Tyr Phe Cys  $_{90}^{\rm MS}$  Ala Arg Tyr Tyr Asp Asp His Tyr Cys Leu Asp Tyr Trp Gly Gln Gly  $_{100}^{\rm MS}$   $_{100}^{\rm MS}$   $_{100}^{\rm MSS}$ 

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 14:

Gin Val Gin Leu Val Gin Ser Gly Gly Car Val Cin Prociety Arg 1 16 15

Ser Leu Arg Leu Ser Cys Lys Ala Ser Gly Tyr Thr Phe Thr Arg Tyr 20 25

Thr Met His Trp Val Arg Gin Ala Pro Gly Lys Gly Leu Glu Trp Ile As 15 Try Val Arg Gin Ala Pro Gly Lys Gly Leu Glu Trp Ile Cys Tyr Ile Asn Pro Ser Arg Gly Tyr Thr Asn Tyr Asn Gln Lys Val 50

Lys Asp Arg Phe Thr Ile Ser Thr Asp Lys Ser Lys Ser Thr Ala Phe 65

Leu Gin Met Asp Ser Leu Arg Pro Glu Asp Thr Ala Val Tyr Tyr Cys 85

Ala Arg Tyr Tyr Asp Asp His Tyr Cys Leu Asp Tyr Trp Gly Gin Gly Thr Thr Leu In Val Ser Ser

(2) INFORMATION FOR SEQ ID NO: 15:

15

- (i) SEQUENCE CHARACTERISTICS:
  (A) LENGTH: 119 amino acids
  (B) TYPE: amino acid
  (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear
  (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEG ID NO: 15:

    Gin Val Gin Leu Val Gin Ser Giy Giy Giy Val Val Val Gin Pro Giy Arg

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    Ser Leu Arg Leu Ser Cys Lys Ala Ser Giy Tyr Thr Phe Thr Arg Tyr

    20

    Thr Het His Trp Val Arg Gin Ala Pro Giy Lys Giy Leu Giu Trp Ile

    45

    Giy Tyr Ile Asn Pro Ser Arg Giy Tyr Thr Asn Tyr Asn Gin Lys Val

    55

    Giy Tyr Tie Asn Pro Ser Arg Giy Tyr Thr Asn Tyr Asn Gin Lys Val

    65

    Lys Asp Arg Phe Thr Ile Ser Thr Asp Lys Ser Lys Ser Thr Ala Phe

    65

    Leu Gin Met Asp Ser Leu Arg Pro Giu Asp Thr Giy Val Tyr Phe Cys

    86

    Ala Arg Tyr Tyr Asp Asp His Tyr Cys Leu Asp Tyr Trp Giy Gin Giy

    Thr Thr Leu Thr Val Ser Ser
- (2) INFORMATION FOR SEQ ID NO: 16:
  - (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 119 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- - (2) INFORMATION FOR SEQ ID NO: 17:
    - (i) SEQUENCE CHARACTERISTICS:
      - (A) LENGTH: 119 amino acids (B) TYPE: amino acid
      - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein

- - Lys Asp arg the first let set and asp asi 75 sys asi in the set of 56 so 70 strain and 176 set of 58 so 175 strain and 176 str

Ala Arg Tyr Tyr Asp Asp His Tyr Cys Leu Asp Tyr Trp Gly Gln Gly
100 105

Thr Thr Leu Thr Val Ser Ser

(2) INFORMATION FOR SEQ ID NO: 18:

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- (i) SEQUENCE CHARACTERISTICS:

  (A) LENGTH: 119 amino acids
  (B) TYPE: amino acid
  - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (x1) SEQUENCE DESCRIPTION: SEQ ID NO: 18:

Gin Val Gin Leu Val Gin Ser Gly Gly Gly Val Val Gin Pro Gly Arg

1 5 10 15 The Pro Gly Arg

Thr Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Ile 35 40

Gly Tyr Ile Asn Pro Ser Arg Gly Tyr Thr Asn Tyr Asn Gln Lys Val 50 55

Lys Asp Arg Phe Thr Ile Ser Thr Asp Lys Ser Lys Asn Thr Leu Phe 65 75 75 86 80 80 Leu Gln Met Asp Ser Leu Arg Pro Glu Asp Thr Gly Val Tyr Phe Cys 95 85 90 90 90

Ala Arg Tyr Tyr Asp Asp His Tyr Cys Leu Asp Tyr Trp Gly Gln Gly  $100\,$ 

Thr Thr Leu Thr Val Ser Ser

- (2) INFORMATION FOR SEQ ID NO: 19:
  - (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 119 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: Single (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 19:

Gln Val Gln Leu Val Gln Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 15

Ser Leu Arg Leu Ser Cys Lys Ala Ser Gly Tyr Thr Phe Thr Arg Tyr 20 25 30

Thr Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Ile 35 40 45

	Gly	Tyr 50	Ile	Asn	Pro	ser	Arg 55	Gly	Tyr	Thr	Asn	Tyr 60	Asn	Gln	Lys	Val
	Lys 65	Asp	Arg	Phe	Thr	Ile 70	Ser	Arg	Asp.	Asn	Ser 75	Lys	Asn	Thr	Leu	Phe 80
	Leu	Gln	Met	Asp	Ser 85	Leu	Arg	Pro	Glu	Asp 90	Thr	Gly	Val	Tyr	Phe 95	Cys
	Ala	Arg	Tyr	Tyr 100	Asp	Asp	His	Tyr	Cys 105	Leu	Asp	Tyr	Trp	Gly 110	Gln	Gly
	Thr	Thr	Leu 115	Thr	Val	Ser	ser									
(2)	INFO	RMAT:	ON I	or :	SEQ :	ID N	0: 2	0:								
	(i)	(A (B	JENCI LEI TYI STI	IGTH PE: RAND	: 11: amin EDNE:	9 am: o ac: SS: :	ino id sing	acid	s							
	(ii)	MOL	ECUL	E TY	PE:	prot	ein									
	(xi)	SEQ	UENC	E DE	SCRI	PTIO	N: S	EQ I	D NO	: 20	:					
	Gln 1	Val	Gln	Leu	Val 5	Gln	Ser	Gly	Gly	Gly 10	Val	Val	Gln	Pro	Gly 15	Arg
	Ser	Leu	Arg	Leu 20	Ser	Cys	Lys	Ala	Ser 25	Gly	Tyr	Thr	Phe	Thr 30	Arg	Tyr
	Thr	Met	His 35	Trp	Val	Arg	Gln	Ala 40	Pro	Gly	Lys	Gly	Leu 45	Glu	Trp	Val
	Ala	Tyr 50	Ile	Asn	Pro	Ser	Arg 55	Gly	Tyr	Thr	Asn	Tyr 60	Asn	Gln	Lys	Phe
	Lys 65	Asp	Arg	Phe	Thr	Ile 70	Ser	Arg	Asp	Asr	75	Lys	Asr	Thr	Leu	Phe 80
	Leu	Gln	Met	Asp	Ser 85	Leu	Arç	Pro	Glu	Asp 90	Thr	Gly	Va]	Tyr	Phe 95	Cys
	Ala	Arg	Tyr	Tyr 100	Asp	Asp	His	туг	Cys 105	Let	) Asp	Tyr	Tr	Gly 110	Gln	Gly
	Thr	Thr	Leu 115		Va)	Ser	Ser									
					200	TO 1										

(2) INFORMATION FOR SEQ ID NO: 21:

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- (i) SEQUENCE CHARACTERISTICS:

  (A) LENGTH: 118 amino acids
  (B) TYPE: amino acid
  (C) STRANDEDMESS: single
  (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 21:

- (2) INFORMATION FOR SEQ ID NO: 22:
  - (i) SEQUENCE CHARACTERISTICS:
    (A) LENGTH: 118 amino acids
    (B) TYPE: amino acid
    (C) STRANDEDNESS: single
    (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 22:

    Gin Val Gin Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 15

    Ser Leu Arg Leu Ser Cys Ser Ala Ser Gly Tyr Thr Phe Thr Arg Tyr 20 25

    Thr Net His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Ile 45 40 40 40 40

    Gly Tyr Ile Asn Pro Ser Arg Gly Tyr Thr Asn Tyr Asn Lys Val Lys 55 50

    Amp Arg Phe Thr Ile Ser Thr Asp Lys Ser Thr Ala Phe Leu 65 67

    Gln Met Asp Ser Leu Arg Pro Glu Asp Thr Ash Val Tyr Tyr Cys Ala 85 100

    Arg Tyr Tyr Amp Amp His Tyr Cys Leu Amp Tyr Trp Gly Gln Gly Thr Leu Thr Val Ser Ser
- (2) INFORMATION FOR SEQ ID NO: 23:

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(i) SEQUENCE CHARACTERISTICS:
 (A) LENGTH: 118 amino acids
 (B) TYPE: amino acid

- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

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- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 23:
- Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg
- Ser Leu Arg Leu Ser Cys Ser Ser Ser Gly Tyr Thr Phe Thr Arg Tyr  $\frac{1}{20}$
- Thr Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Ile 35 40 45
- Gly Tyr Ile Asn Pro Ser Arg Gly Tyr Thr Asn Tyr Asn Lys Val Lys 50 60
- Asp Arg Phe Thr Ile Ser Thr Asp Lys Ser Lys Ser Thr Ala Phe Leu 65 70 75 80
- Gln Met Asp Ser Leu Arg Pro Glu Asp Thr Ala Val Tyr Tyr Cys Ala 85 90 95
- Arg Tyr Tyr Asp Asp His Tyr Cys Leu Asp Tyr Trp Gly Gln Gly Thr
- Thr Leu Thr Val Ser Ser
- (2) INFORMATION FOR SEQ ID NO: 24:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 118 amino acids
      (B) TYPE: amino acid
      (C) STRANDEDNESS: single
      (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 24:
- Gln Val Gln Leu Val Gln Ser Gly Gly Gly Val Val Gln Pro Gly Arg
- Ser Leu Arg Leu Ser Cys Ser Ala Ser Gly Tyr Thr Phe Thr Arg Tyr 20 25 30
- Thr Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Ile 35 40 45
- Gly Tyr Ile Asn Pro Ser Arg Gly Tyr Thr Asn Tyr Asn Lys Val Lys 50 60Asp Arg Phe Thr Ile Ser Thr Asp Lys Ser Lys Ser Thr Ala Phe Leu
- Gln Met Asp Ser Leu Arg Pro Glu Asp Thr Gly Val Tyr Phe Cys Ala 85 90 95
- Arg Tyr Tyr Asp Asp His Tyr Cys Leu Asp Tyr Trp Gly Gln Gly Thr

Thr Leu Thr Val Ser Ser 115

- (2) INFORMATION FOR SEQ ID NO: 25:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 118 amino acids (B) TYPE: amino acid

      - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein

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- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 25:
- Gln Val Gln Leu Val Glu Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 10 15
- Ser Leu Arg Leu Ser Cys Ser Ala Ser Gly Tyr Thr Phe Thr Arg Tyr  $\frac{1}{20}$
- Thr Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Ile 35 40 45
  - Gly Tyr Ile Asn Pro Ser Arg Gly Tyr Thr Asn Tyr Asn Lys Val Lys 50 60
    - Asp Arg Phe Thr Ile Ser Thr Asp Lys Ser Lys Ser Thr Ala Phe Leu 65 70 75 80
    - Gln Met Asp Ser Leu Arg Pro Glu Asp Thr Gly Val Tyr Phe Cys Ala as 90 95
    - Arg Tyr Tyr Asp Asp His Tyr Cys Leu Asp Tyr Trp Gly Gln Gly Thr 100 105
  - Thr Leu Thr Val Ser Ser 115
- (2) INFORMATION FOR SEQ ID NO: 26:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 118 amino acids (B) TYPE: amino acid
    - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein
- - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 26:
  - Gln Val Gln Leu Val Gln Ser Gly Gly Gly Val Val Gln Pro Gly Arg 1 5 10 15
- Ser Leu Arg Leu Ser Cys Ser Ala Ser Gly Tyr Thr Phe Thr Arg Tyr 25 30
- Thr Met His Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Ile 35 40 45
- 50 Gly Tyr Ile Asn Pro Ser Arg Gly Tyr Thr Asn Tyr Asn Lys Val Lys 50 60

		ASP 65	Arg	Pne	THE	Tie	70	1111	мэр	2,0		75	,	ŧ.			80
		Gln	Met	Asp	Ser	Leu 85	Arg	Pro	Glu	Asp	Th: 90	Gly	Val	Tyr	Phe	Cys 95	Ala
5		Arg	Tyr	Tyr	Asp 100	Asp	His	туг	Cys	Leu 105	Asp	Tyr	Trp	Gly	Gln 110	Gly	Thr
		Thr	Leu	Thr 115	Val	ser	ser										
10	(2)	INFO			FOR S	SEQ :	ED N	): 2	7:								
15		(i)	(A (B (C	) LE	E CHA NGTH PE: A RANDA POLO	: 12 min EDNE:	sam: sac:	ino : id sing	acid:	5							
		(ii)	MOL	ECUL	E TY	PE: 1	prot	ein									
20		(xi)	SEQ	UENC	E DE	SCRI	PTIO	N: S	EQ I	D NO	: 27	:					
		Gln 1	Val	Gln	Leu	Val	Glu	Ser	Gly	Gly	Gly 10	Val	Va1	Gln	Pro	Gly 15	Arg
ne.		Ser	Leu	Arg	Leu 20	Ser	Cys	Ser	Ser	Ser 25	Gly	Phe	Ile	Phe	Ser 30	Ser	Tyr
25		λla	Met	Tyr 35	Trp	Val	Arg	Gln	Ala 40	Pro	Gly	Lys	Gly	Leu 45	Glu	Trp	Val
		Ala	Ile 50	Ile	Trp	Asp	Asp	G1y 55	Ser	Asp	Gln	His	TY1	Ala	Asp	Ser	Val
30		Lys 65	Gly	Arg	Phe	Thr	11e 70	Ser	Arg	Asp	Asn	Ser 75	Lys	Asn	Thr	Leu	Phe 80
		Leu	Glr	Met	: Asp	Set 85	Leu	Arç	Pro	Glu	Asp 90	Thr	Gly	Val	Tyr	Phe 95	Cys
35		Ala	Arç	j Asp	Gly	Gly	His	Gly	Phe	Cys	Ser	Ser	Ala	ser	Cys	Phe	Gly
		Pro	Asp	Ty:	r Trp	Gl <sub>y</sub>	Glr	Gly	Thr 120	Pro	Val	The	· Va	125	Ser		
40	(2)	INFO	RMA'	CION	FOR	SEQ	ID N	10: 3	28:								
		(i)	(1	A) LI B) T	CE CI ENGTI YPE: TRANI	i: 10 ami: DEDNI	)7 ar 10 ac ESS:	nino cid sin	ació	ls							
45		(ii			OPOLO												
50					CE D								t Se	r Al	a Sei	Pro	Gly
		1				5					10					15	
55																	

	Glu	Lys	Val	Thr 20	Met	Thr	Cys	Ser	Ala 25	Ser	Ser	Ser	Val	Ser 30	Tyr	Met
	Asn	Trp	Tyr 35	Gln	Gln	Lys	Ser	Gly 40	Thr	Ser	Pro	Lys	A∠g 45	Trp	Ile	Tyr
	Asp	Thr 50	Ser	Lys	Leu	Ala	Ser 55	Gly	Val	Pro	Ala	His 60	Phe	Arg	Gly	Ser
	Gly 65	Ser	Gly	Thr	Ser	Tyr 70	Ser	Leu	Thr	Ile	Ser 75	Gly	Met	Glu	Ala	Glu 80
	Asp	Ala	Ala	Thr	Tyr 85	Tyr	Cys	Gln	Gln	Trp 90	Ser	Ser	Asn	Pro	Phe 95	Thr
	Phe	Gly	Ser	Gly 100	Thr	Lys	Leu	Glu	Ile 105	Asn	Arg					
(2)	INFO	RMAT	ION	FOR	SEQ	ID N	0; 2	9:								
	(i)	(A (B (C	) LE ) TY ) ST	ngth PE: Rand	ARAC : 10 amin EDNE GY:	7 am o ac SS:	ino id sing	acid	5							
	(ii)	MOL	ECUL	Е ТУ	PE:	prot	ein									
	(xi)	SEQ	UENC	E DE	SCRI	PTIO	N: S	EQ I	р ио	: 29	:					
	Asp 1	Ile	Gln	Met	Thr 5	Gln	Ser	Pro	Ser	Ser 10	Leu	Ser	Ala	Ser	Val 15	Gly
	Asp	Arg	Va 1	Thr 20	Ile	Thr	Cys	Ser	Ala 25	Ser	Ser	Ser	Val	Ser 30	Tyr	Met
			35					40					45	Leu		
	Asp	Thr 50	Ser	Lys	Leu	Ala	Ser 55	Gly	Val	Pro	Ser	Arg 60	Phe	Ser	Gly	Ser
	65					70					75					Glu 80
	Asp	Ile	: Ala	Thi	Ty1	Tyr	- Cys	Glr	Gln	Trp 90	Ser	Ser	Asr	Pro	Phe 95	Thr
	Phe	61)	Gli	100	Th:	Lys	Leu	Glr	105	Thr	Arc	•				
(2)	INFO	ORMAT	NOI	FOR	SEQ	ID I	10: 3	:0								
	(i)	(1	A) L: 3) T: C) S'	ENGT (PE: [RAN]	HARAG H: 16 amin DEDNI DGY:	07 aı no ad ESS:	aino cid sino	acio	ls							
	(ii	MO:	LECU	LE T	YPE:	pro	tein									
	(xi	) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ :	ED NO	): 30	):					

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(2) INFORMATION FOR SEQ ID NO: 31:

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- (i) SEQUENCE CHARACTERISTICS:

  (A) LENGTH: 107 amino acids
  (B) TYPE: amino acid
  (C) STRANDEDNESS: single
  (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ IO NO: 31:

  Gln Ile Val Mat Thr Gln Ser Pro Ser Ser Leu Ser Ala Ser Val Gly 15

  Asp Arg Val Thr Ile Thr Cys Ser Ala Ser Ser Ser Val Ser Tyr Met 25

  Asn Trp Tyr Gln Gln Thr Pro Gly Lys Ala Pro Lys Arg Trp Ile Tyr 45

  Asp Thr Ser Lys Leu Ala Ser Gly Val Pro Ser Arg Phe Ser Gly Ser 55

  Gly Ser Gly Thr Asp Tyr Thr Phe Thr Ile Ser Ser Leu Gln Pro Glu 75

  Asp Ile Ala Thr Tyr Cys Gln Gln Trp Ser Ser Asn Pro Phe Thr 85

  Phe Gly Gln Gln Tyr Lys Leu Gln Ile Thr Arg 105
- (2) INFORMATION FOR SEQ ID NO: 32:
  - (i) SEQUENCE CHARACTERISTICS:

    (A) LENGTH: 107 amino acids
    (B) TYPE: amino acid
    (C) STRANDEDNESS: single
    (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 32: Asp Ile Gln Met Thr Gln Ser Pro Ser Ser Leu Sariala Ser Wel Gly Asp Arg Val Thr Ile Thr Cys Ser Ala Ser Ser Ser Val Ser Tyr Met Asn Trp Tyr Gln Gln Thr Pro Gly Lys Ala Pro Lys Arg Trp Tle Tyr Asp Thr Ser Lys Leu Ala Ser Gly Val Pro Ser Arg Phe Ser Gly Ser Gly Ser Gly Thr Asp Tyr Thr Phe Thr Ile Ser Ser Leu Gln Pro Glu Asp Ile Ala Thr Tyr Tyr Cys Gln Gln Trp Ser Ser Asn Pro Phe Thr 85 90 95 Phe Gly Gln Gly Thr Lys Leu Gln Ile Thr Arg (2) INFORMATION FOR SEQ ID NO: 33: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 108 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: protein (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 33: Asp Ile Gln Met Thr Gln Ser Pro Ser Ser Leu Ser Ala Ser Val Gly Asp Arg Val Thr Ile Thr Cys Gln Ala Ser Gln Asp Ile Ile Lys Tyr

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Glu Asp Ile Ala Thr Tyr Tyr Cys Gln Gln Tyr Gln Ser Leu Pro Tyr 95 Thr Phe Gly Gln Gly Thr Lys Leu Gln Ile Thr Arg

Leu Asn Trp Tyr Gln Gln Thr Pro Gly Lys Ala Pro Lys Leu Leu Ile  $_{35}$ Tyr Glu Ala Ser Asn Leu Gln Ala Gly Val Pro Ser Arg Phe Ser Gly Ser Gly Ser Gly Thr Asp Tyr Thr Phe Thr Ile Ser Ser Leu Gln Pro

Thr Phe Gly Gln Gly Thr Lys Leu Gln Ile Thr Ary 100 105

# Claims

 A CDR-grafted antibody heavy chain having a variable region domain comprising acceptor framework and donor antigen binding regions wherein the framework comprises donor residues at at least one of positions 6, 23 and/or 24, 48 and/or 49, 71 and/or 73, 73 and/or 78 and/or 78 and/or 80 and/or 91.

- 2. A CDR-grafted heavy chain according to Claim 1 comprising donor residues at positions 23, 24, 49, 71, 73 and 78, or at positions 23, 24 and 49.
- 3. A CDR-grafted heavy chain according to Claim 2 comprising donor residues at positions 2, 4, 6, 25, 36, 37, 39, 47, 48, 93, 94, 103, 104, 106 and 107.
  - 4. A CDR-grafted heavy chain according to Claim 2 or 3, comprising donor residues at one, some or all of positions:
  - 1 and 3.
  - 69 (if 48 is different between donor and acceptor),
    - 38 and 46 (if 48 is the donor residue),

    - 82 and 18 (if 67 is the donor residue).

- any one or more of 9, 11, 41, 87, 108, 110 and 112.
  - 5. A CDR-grafted heavy chain according to any of the preceding comprising donor CDRs at positions 26-35, 50-65 and 95-100.
- 20 6. A CDR-grafted antibody light chain having a variable region domain comprising acceptor framework and donor antigen binding regions wherein the framework comprises donor residues at at least one of positions 1 and/or 3 and 46 and/or 47.
  - A CDR-grafted light chain according to Claim 6 comprising donor residues at positions 46 and 47.
  - 8. A CDR-grafted antibody light chain having a variable region domain comprising acceptor framework and donor antigen binding regions wherein the framework comprises donor residues at at least one of positions 46, 48, 58 and 71.
- 30 9. A CDR-grafted light chain according to Claim 8 comprising donor residues at positions 46, 48, 58 and 71.
  - 10. A CDR-grafted light chain according to Claim 8 or 9, comprising donor residues at positions 2, 4, 6, 35, 36, 38, 44, 47, 49, 62, 64-69, 85, 87, 98, 99, 101 and 102.
- 35 11. A CDR-grafted light chain according to Claim 9 or 10, comprising donor residues at one, some or all of positions: 1 and 3.
  - 63.
  - 60 (if 60 and 54 are able to form a potential saltbridge),
    - 70 (if 70 and 24 are able to form a potential saltbridge),
      - 73 and 21 (if 47 is different between donor and acceptor),
      - 37 and 45 (if 47 if different between donor and acceptor), and

    - any one or more of 10, 12, 40, 83, 103 and 105.
  - 12. A CDR-grafted light chain according to any one of Claims 6-11, comprising donor CDRs at positions 24-34, 50-56 and 89-97.
  - 13. A CDR-grafted antibody molecule comprising at least one CDR-grafted heavy chain according to any one of Claims 1-5 and at least one CDR-grafted light chain according to any one of Claims 6-12.
    - 14. A CDR-grafted antibody molecule according to Claim 13, which is a site-specific antibody molecule.
  - 15. A CDR-grafted antibody molecule according to Claim 13 which has specificity for an interleukin, hormone or other biologically active compound or a receptor therefor.
  - 16. A CDR-grafted antibody heavy or light chain or molecule according to any one of the preceding claims comprising human acceptor residues and non-human donor residues.

- A DNA sequence which codes for a CDR-grafted heavy chain according to Claim 1 or a CDR-grafted light chain according to Claim 6 or Claim 8.
- 18. A cloning or expression vector containing a DNA sequence according to Claim 17.
- 19. A host cell transformed with a DNA sequence according to Claim 17.

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- 20. A process for the production of a CDR-grafted antibody sequence according to Claim 17 in a transformed host cell.
- 21. A process for producing a CDR-grafted antibody product comprising: (a) producing in an expression vector an operon having a DNA sequence which encodes an antibody heavy chain according to Claim 1;
- and/or (b) producing in an expression vector an operon having a DNA sequence which encodes a complementary antibody light chain according to Claim 6 or Claim 8: (c) transferding a host coll with the or each vector;
  - and
     (d) culturing the transfected cell line to produce the CDR-grafted antibody product.
- 22. A therapeutic or diagnostic composition comprising a CDR-grafted antibody heavy chain according to Claim 1, or a CDR-grafted light chain according to Claim 6 or Claim 8, or a CDR-grafted antibody molecule according to Claim 13 in combination with a pharmaceutically acceptable carrier, diluent or excipient.
  - 23. A method of therapy or diagnosis comprising administering an effective amount of a CDR-grafted heavy chain according to Claim 1, or a CDR-grafted light chain according to Claim 6 or Claim 6, or a CDRgrafted artibody molecule according to Galim 13 to a human or animal subject.

- 1 GAATTCCCAA AGACAAAatg gattttcaag tgcagatttt cagcttcctg 51 ctaatcagtg cctcagtcat aatatccaga ggacaaattg ttctcaccca 101 gtctccagca atcatgtctg catctccagg ggagaaggtc accatgacct 151 gcaqtqccaq ctcaaqtqta aqttacatqa actqqtacca qcaqaaqtca 201 ggcacctccc ccaaaagatg gatttatgac acatccaaac tggcttctgg .251 agtccctqct cacttcaggg qcaqtggqtc tgggacctct tactctctca 301 caatcagcgg catggaggct gaagatgctg ccacttatta ctgccagcag 351 tggagtagta acccattcac gttcggctcg gggacaaagt tggaaataaa 401 ccgggctgat actgcaccaa ctgtatccat cttcccacca tccagtgagc 451 agttaacatc tqqaqqtqcc tcaqtcqtqt gcttcttqaa caacttctac 501 cccaaagaca tcaatgtcaa gtggaagatt gatggcagtg aacgacaaaa 551 tggcqtcctg aacagttgga ctgatcagga cagcaaagac agcacctaca 601 gcatgagcag cacceteacg ttgaccaagg acgagtatga acgacataac 651 agctatacct gtgaggccac tcacaagaca tcaacttcac ccattgtcaa 701 gagetteaac aggaatgagt gtTAGAGACA AAGGTCCTGA GACGCCACCA 751 CCAGCTCCCA GCTCCATCCT.ATCTTCCCTT CTAAGGTCTT GGAGGCTTCC 801 CCACAAGCGC tTACCACTGT TGCGGTGCTC tAAACCTCCT CCCACCTCCT 851 TCTCCTCCTC CTCCCTTTCC TTGGCTTTTA TCATGCTAAT ATTTGCAGAA 901 AATATTCAAT AAAGTGAGTC TTTGCCTTGA AAAAAAAAA AAA
  - Fig. 1(a)
  - 1 MDFOVOIFSF LLISASVIIS RGQIVLTQSP AIMSASPGEK VTMTCSASSS
  - 51 VSYMNWYQQK SGTSPKRWIY DTSKLASGVP AHFRGSGSGT SYSLTISGME
  - 101 AEDAATYYCQ QWSSNPFTFG SGTKLEINRA DTAPTVSIFP PSSEQLTSGG
  - 151 ASVVCFLNNF YPKDINVKWK IDGSERONGV LNSWTDODSK DSTYSMSSTI.
- 201 TIJTKDEYERH NSYTCEATHK TSTSPIVKSF NRNEC\*

Fig. 1(b)

1	GAATTCCCCT CTCCACAGAC ACTGAAAACT CTGACTCAAC ATGGAAAGGC
51	ACTGGATCTT TCTACTCCTG TTGTCAGTAA CTGCAGGTGT CCACTCCCAG
101	GTCCAGCTGC AGCAGTCTGG GGCTGAACTG GCAAGACCTG GGGCCTCAGT
151	GAAGATGTCC TGCAAGGCTT CTGGCTACAC CTTTACTAGG TACACGATGC
201	ACTGGGTAAA ACAGAGGCCT GGACAGGGTC TGGAATGGAT TGGATACATT
251	AATCCTAGCC GTGGTTATAC TAATTACAAT CAGAAGTTCA AGGACAAGGC
301	CACATTGACT ACAGACAAAT CCTCCAGCAC AGCCTACATG CAACTGAGCA
351	GCCTGACATC TGAGGACTCT GCAGTCTATT ACTGTGCAAG ATATTATGAT
401	GATCATTACT GCCTTGACTA CTGGGGCCAA GGCACCACTC TCACAGTCTC
451	CTCAGCCAAA ACAACAGCCC CATCGGTCTA TCCACTGGCC CCTGTGTGTG
501	GAGATACAAC TGGCTCCTCG GTGACTCTAG GATGCCTGGT CAAGGGTTAT
551	TTCCCTGAGC CAGTGACCTT GACCTGGAAC TCTGGATCCC TGTCCAGTGG
601	TGTGCACACC TTCCCAGCTG TCCTGCAGTC TGACCTCTAC ACCCTCAGCA
651	GCTCAGTGAC TGTAACCTCG AGCACCTGGC CCAGCCAGTC CATCACCTGC
701	AATGTGGCCC ACCCGGCAAG CAGCACCAAG GTGGACAAGA AAATTGAGCC
751	CAGAGGGCCC ACAATCAAGC CCTGTCCTCC ATGCAAATGC CCAGCACCTA
801	ACCTCTTGGG TGGACCATCC GTCTTCATCT TCCCTCCAAA GATCAAGGAT
851	GTACTCATGA TCTCCCTGAG CCCCATAGTC ACATGTGTGG TGGTGGATGT
901	GAGCGAGGAT GACCCAGATG TCCAGATCAG CTGGTTTGTG AACAACGTGG
951	AAGTACACAC AGCTCAGACA CAAACCCATA GAGAGGATTA CAACAGTACT
1001	CTCCGGGTGG TCAGTGCCCT CCCCATCCAG CACCAGGACT GGATGAGTGG
1051	CANGGAGTTC AAATGCAAGG TCAACAACAA AGACCTCCCA GCGCCCATCG
1101	AGAGAACCAT CTCAAAACCC AAAGGGTCAG TAAGAGCTCC ACAGGTATAT
1151	GTCTTGCCTC CACCAGAAGA AGAGATGACT AAGAAACAGG TCACTCTGAC
1201	CTGCATGGTC ACAGACTTCA TGCCTGAAGA CATTTACGTG GAGTGGACCA
1251	ACAACGGGAA AACAGAGCTA AACTACAAGA ACACTGAACC AGTCCTGGAC
1301	
1351	
1401	
1451	
1501	TGCTTCCCTT GTATAAATAA AGCACCCAGC AATGCCTGGG ACCATGTAAA

Fig. 2(a)

OKT 3 HEAVY CHAIN PROTEIN SEQUENCE DEDUCED FROM DNA SEQUENCE

```
1 MERHWIFLLL LSVTAGVHSQ VQLQQSGAEL ARPGASVKMS CKASGYTFTR
51 YTMHWVKQRP GQGLEWIGYI NPSRGYTNYN QKFKDKATLT TDKSSSTAYM
101 QLSSLTSEDS AVYYCARYYD DHYCLDYWGQ GTTLTVSSAK TTAPSVYPLA
151 PVCGDTTGSS VTLGCLVKGY FPEPVTLTWN SGSLSSGVHT FPAVLQSDLY
201 TLSSSVTVTS STWPSQSITC NVAHPASSTK VDKKIEPRGP TIKPCPPCKC
251 PAPNILICOPS UFTEPPKIKD VIMISLSPIV TCVVVDVSED DPDVQISWFV
301 NNVEVHTAQT QTHREDYNST LRVVSALPIQ HODWMSGKEF KCKVNNKDLP
351 APIERTISKP KGSVRAPQVY VLPPPEEEMT KKQVTLTCMV TDFMPEDIYV
401 EWTNNGKTEL NYKNTEPVLD SDGSYFMYSK LRVEKKNWVE RNSYSCSVVH
451 EGLHNHHTTK SFSRTPGK*
                                 Fig. 2(b)
                                23
                                                    42
           1
           NN
                   N
           SBspSPESssBSbSsSssPSPSPsPSsse*s*p*Pi^ISsSe
RES TYPE
           OIVLTOSPAIMSASPGEKVTMTCSASS.SVSYMNWYQQKSGT
Okt3vl
REI
           DIQMTQSPSSLSASVGDRVTITCQASQDIIKYLNWYQQTPGK
           ? ?
             CDR1
                   (LOOP)
             CDR1
                   (KABAT)
                                                    85
                       56
          N NN
RES TYPE *IsiPpIeesesssSBEsePsPSBSSEsPspsPsseesSPePb
          SPKRWIYDTSKLASGVPAHFRGSGSGTSYSLTISGMEAEDAAT
Okt3v1
          APKLLIYEASNLQAGVPSRFSGSGSGTDYTFTISSLQPEDIAT
RET
           7 77
                 ****** CDR2 (LOOP/KABAT)
                         102
RES TYPE PiPIPies**iPIIsPPSPSPSS
                                            Fig. 3
          YYCQQWSSNPFTFG8GTKLEINR
Okt3vl
         YYCQQYQSLPYTFGQGTK<u>LQ</u>I<u>T</u>R
REIvl
                              2
                            CDR3 (LOOP)
                            CRD3 (KABAT)
```

	NN N		23 26	32 35	N39 43	
RES TYPE	SESPs SBssS	sSSsSpSpS	PsPSEbSI	BssBePi	PIpiesss	
Okt3h	QVQLQQSGAEL					
KOL	OVOLVESGGGY					
	?		??			
			*:	****	CDR1 (L	00P)
				****	* CDR1 (KA	BAT)
					•	
	52a	60	65 N	N N	82abc	89
RES TYPE I	Telppp sssss	sss <sup>*</sup> ps <sup>*</sup> pS	SsbSpseS	sSseSp^	pSpsSBssS	^ePb
	LEWIGYINPSRG					
	LEWVALIWDDGS					
	??		? ?	? ?		?
	*****	*****	CD	R2 (LC	OP)	
	*****	******	**** CD	R2 (KA	BAT)	
	92 N		107	i13		
RES TYPE	PiPIEissssi	iisssbibi	*EIPIP*s	pSBSS		
Okt3vh	YYCARYYDDHY	cL	DYWGQGTT	LTVSS		
KOL	YFCARDGGHGF	CSSASCFGP	DYWGQGTI	VTVSS		
		******			(LOOP)	

Fig. 4

# OKT 3 HEAVY CHAIN CDR GRAFTS

# 1. gh341 and derivatives

	1 26 35 39 43	
kt3vh	QVQLQQSGAELARPGASVKMSCKASGYTFTRYTMHWVKQRPGQ	
H341	OVOLVESGGGVVQPGRSLRLSCSSSGYTFTRYTMHWVRQAPGK	JA178
H341A	QVQLVQSGGGVVQPGRSLRLSC <u>KASGYTFTRYTM</u> HWVRQAPGK	JA185
gH341E	QVQLVQSGGGVVQPGRSLRLSC <u>KASGYTFTRYTM</u> HWVRQAPGK	JA198
gH341*	OVOLVOSGGGVVQPGRSLRLSCKASGYTFTRYTMHWVRQAPGK	JA207
gH341*	OVOLVOSGGGVVQPGRSLRLSCKASGYTFTRYTMHWVRQAPGK	JA209
gH341D	OVOLVOSGGGVVQPGRSLRLSC <u>KASGYTFTRYTM</u> HWVRQAPGK	JA197
gH341*	OVOLVOSGGGVVQPGRSLRLSC <u>KASGYTFTRYTM</u> HWVRQAPGK	JA199
gH341C	QVQLVQSGGVVQPGRSLRLSC <u>KASGYTFTRYTM</u> HWVRQAPGK	JA184
gH341*	QVQLVQSGGGVVQPGRSLRLSCS <u>ASGYTFTRYTM</u> HWVRQAPGK	JA203
gH341*	QVQLVESGGGVVQPGRSLRLSCS <u>ASGYTFTRYTM</u> HWVRQAPGK	JA205
gH341B	QVQLVESGGGVVQPGRSLRLSCSS <u>SGYTFTRYTM</u> HWVRQAPGK	JA183
gH341*	OVOLVQSGGGVVQPGRSLRLSCSASGYTFTRYTMHWVRQAPGK	JA204
gH341*	OVOLVESGGGVVQPGRSLRLSCSASGYTFTRYTMHWVRQAPGK	JA206
gH341*	QVQLVQSGGGVVQPGRSLRLSCSASGYTFTRYTHHWVRQAPGK	JA208
KOL	QVQLVESGGGVVQPGRSLRLSCSSSGFIFSSYAMYWVRQAPG	<

Fig. 5(i)

	44	50	65	83	
okt3vh	GLE	WIGYINP	srgytnynqkfkDkatli	TDKSSSTAYMQLSSLT	
gH341	GLE	WVAYINP	SRGYTNYNOKFKDRFTI:	SRDNSKNTLFLQMDSLR	JA178
gH341A	GLE	W <u>IGYIN</u> P	<u>SRGYTNYNOKVKD</u> RFTI	S <u>T</u> D <u>K</u> SK <b>S</b> TAFLOMDSLR	JA185
gH341E	GLE	WIGYIND	<u>FRGYTNYNOK</u> V <u>KD</u> RFT15	TDKSKSTAFLQMDSLR	JA198
gH341*	GLE	WIGYINP:	<u>SRGYTNYNOKVKD</u> RFTIS	TDKSKNTAFLQMDSLR	JA207
gH341*	GLE	WIGYINE	<u>SRGYTNYNOK</u> V <u>KD</u> RFTIS	RDNSKNT <u>A</u> FLQMDSLR	JA209
gH341D	GLE	WIGYINP:	<u>SRGYTNYNOKVKD</u> RFTIS	<u>TDK</u> SKNTLFLQMDSLR	JA197
gH341*	GLE	WIGYINP	<u>SRGYTNYNOK</u> V <u>KD</u> RFTIS	RDNSKNTLFLQMDSLR	JA199
gH341C	GLE	WVA <u>YINP</u>	SRGYTNYNOKFKDRFTIS	RDNSKNTLFLQMDSLR	JA184
gH341*	GLE	WIGYINP	<u>SRGYTNYNOK</u> V <u>KD</u> RFTI	<u>TDKSKSTAFLQMDSLR</u>	JA207
gH341*	GLE	WIGYINP	<u>SRGYTNYNOK</u> V <u>KD</u> RFTI:	TDKSKSTAFLQMDSLR	JA205
gH341B	GLE	WIGYINP	<u>SRGYTNYNOK</u> V <u>KD</u> RFTI:	STOKSKSTAFLQMDSLR	JA183
gH341*	GLE	WIGYINP	SRGYTNYNOKVKDRFTI:	STOKSKSTAFLQMDSLR	JA204
gH341*	GLE	WIGYINP	<u>SRGYTNYNOK</u> V <u>KD</u> RFTI:	STDKSKSTAFLQMDSLR	JA206
gH341*	GLE	WIGYINP	SRGYTNYNOKVKDRFTI:	STOKSKNTAFLQMDSLR	JA208
KOL	GLE	WVAIIWE	DGSDQHYADSVKGRFTI	SRDNSKNTLFLQMDSLR	

Fig. 5(ii)

	84	95	102	113	
Okt3vh	SEDS	AVYYCARYYDDHY.	CLDYWGQG	TTLTVSS	
gH341	PEDT	GVYFCARYYDDHY.	CLDYWGQG	TTLTVSS	JA178
gH341A	PEDT	AVYYCARYYDDHY.	CLDYWGQG	TTLTVSS	JA185
gH341E	PEDT	GVYFCAR <u>YYDDHY</u> .	CLDYWGQG	TTLTVSS	JA198
gH341*	PEDT	GVYFCARYYDDHY.	CLDYWGQG	TTLTVSS	JA207
gH341D	PEDT	GVYFCAR <u>YYDDHY.</u>	CLDYWGQG	TTLTVSS	JA197
gH341*	PEDI	GVYFCARYYDDHY.	CLDYWGQG	TTLTVSS	JA209
gH341*	PEDT	GVYFCARYYDDHY.	CLDYWGQG	TTLTVSS	JA199
gH341C	PEDT	GVYFCAR <u>YYDDHY.</u>	CLDYWGQG	TTLTVSS	JA184
gH341*	PEDT	<u>AVYY</u> CARY <u>YDDHY.</u>	CLDYWGQG	TTLTVSS	JA203
gH341*	PEDI	AVYYCARYYDDHY.	CLDYWGQG	TTLTVSS	JA205
gH341B	PEDT	AVYYCARYYDDHY.	CLDYWGQG	TTLTVSS	JA183
gH341*	PEDI	GVYFCARYYDDHY.	CLDYWGQG	TTLTVSS	JA204
gH341*	PEDI	GVYFCARYYDDHY.	CLDYWGQG	TTLTVSS	JA206
gH341*	PEDT	GVYFCARYYDDHY.	CLDYWGQG	TTLTVSS	JA208
KOL	PED1	GVYFCARDGGHGFC	SSASCFGPDYWGQG	TPVTVSS	

Fig. 5 (iii)

# OKT3 LIGHT CHAIN CDR GRAFTING

# 1. gL221 and derivatives

	1				24	34	42
Okt3vl	QIV	LTQSP	AIMS	ASPGE	KVTMTCSASS.	SVSYMNWYQQ	KSGT
gL221	DIQ	MTQSP	SSLS.	ASVGD	RVTITCSASS.	SVSYMMWYQQ	TPGK
gL221A	OIA	MTQSP	SSLS	ASVGD	RVTITC <u>SASS</u> .	SVSYMNWYQQ	TPGK
gL221B	<u>QIV</u>	MTQSP	SSLS	<b>ASVGD</b>	RVTITC <u>SASS</u> .	SVSYMNWYQQ	TPGK
gL221C	DIQ	MTQSP	SS <b>LS</b>	<b>ASVGD</b>	RVTITC <u>SASS</u> .	SVSYMNWYQC	TPGK
REI	DIC	MTQSF	SSLS	ASVGD	RVTITCQASQI	I I KYLNWYQQ	TPGK
	43	5	0	56			85
Okt3vl					AHFRGSGSGTS		
gL221					SRFSGSGSGT		
gL221A					SRFSGSGSGTI		
gL221B					SRFSGSGSGT		
gL221C					SRFSGSGSGT		
REI	API	KLLIYI	EASNI	QAGVE	ERFSGSGSGT	DYTFTISSLOI	PEDIAT
	86	91	96		108		
Okt3v1	YYC	QWSS	NPFTI	GSGTI	KLEINR		
gL221	YYC	OOWSS	NPFTI	GOGTI	KLQITR		
gL221A	YYC	DOWSS	NPFT	FGQGTI	KLQITR		
gL221B	YYC	oowss	NPFT	FGQGTI	KLQITR		
gL221C	YYC	oowss	NPFT	FGQGTI	KLQITR		

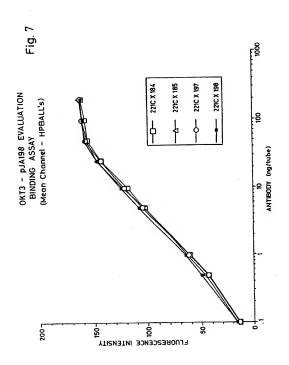
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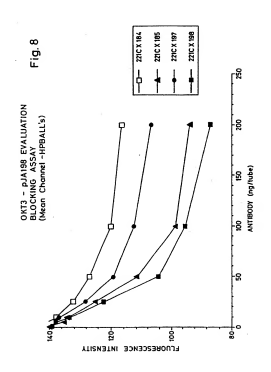
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FRAMEWORK RESIDUES INCLUDED IN THE GENE ARE DOUBLE UNDERLINED

Fig. 6

YYCQQYQSLPYTFGQGTKLQITR





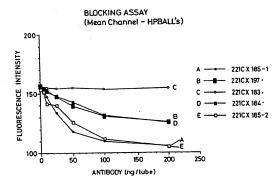
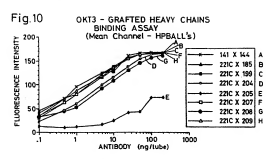
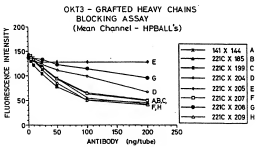
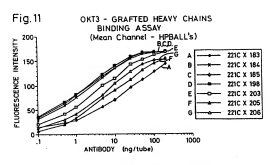


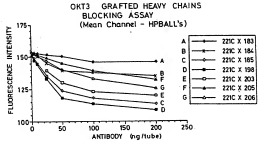
Fig.9



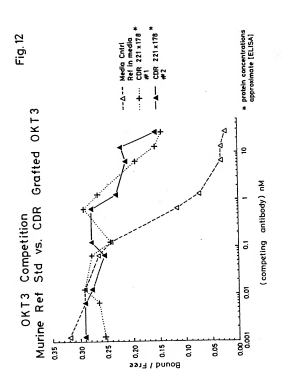


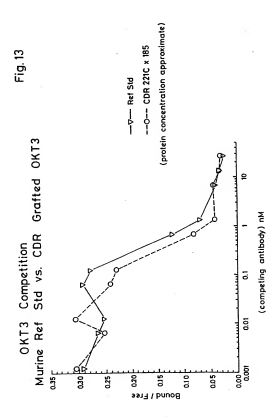
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	141 X	144





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(206)	,24,48,49,71,73,76,78,
—o— (203)	6 , 24,48,49,71,73,76,78,88,91,
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(198)	6.23.24.48.49.71.73.76.78,_







# PARTIAL EUROPEAN SEARCH REPORT

pelication Number

which under Rule 45 of the European Patent Convention EP 94 10 4042 shall be considered, for the purposes of subsequent proceedings, as the European search report

Category D,X	Citation of document with ind of relevant pass PROCEEDINGS OF THE N SCIENCES OF THE USA	ages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.CLS)
D,X		ATIONAL ACADEMY OF	1-22	010015 (12
	vol. 86, no. 24 , 15 WASHINGTON, DC, USA pages 10029 - 10033	December 1989 , umanized antibody that ukin 2 receptor.'		C12M15/13 C07K15/28 A61K39/395 G01N33/577
P,X	EP-A-0 403 156 (GENZ BEHRINGWERKE) * the whole document		1-23	
				TECHNICAL FIELDS SEARCHED (BA.CLS) C12N C07K A61K G01N
The Seat the provi out a me Claims s Claims s	OMPLETE SEARCH  rch Dirichis considers that the present it closes of the Extraoral rabust Covered analogical search size the state of the art sarched completely a rearched completely are arched completely to the streamful of the search in t	Zeropeaa patent application does not compa n to such an extent that it is not possible to on the balls of ones of the dates	y xidb o carry	
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	Place of courch	Date of completion of the search		Examiner
	THE HAGUE	8 June 1994	No	oij, F
	CATEGORY OF CITED DOCUME	VIS I : theory or princi E : earlier patent d after the filing	ple underlying to	he invention blished on, or



EP 94 10 4042

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Remark: Although claim 23
is directed to a method of
treatment of (diagnostic method
practised on) the human/animal body
(Art. 52(4) ERC) the search has been
carried out and based on the
alleged effects of the compound/
composition